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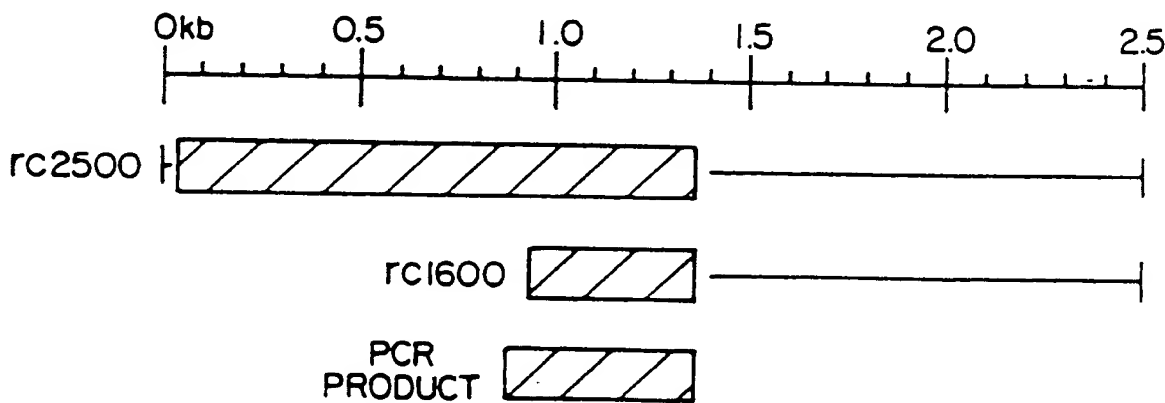
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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5 :  C12N 15/54, 9/10		A3	(11) International Publication Number: WO 92/09694  (43) International Publication Date: 11 June 1992 (11.06.92)
<p>(21) International Application Number: PCT/CA91/00417</p> <p>(22) International Filing Date: 29 November 1991 (29.11.91)</p> <p>(30) Priority data: 620,098 30 November 1990 (30.11.90) US</p> <p>(71) Applicant: HSC RESEARCH AND DEVELOPMENT LIMITED PARTNERSHIP [CA/CA]; 88 Elm Street, Toronto, Ontario M5G 1X8 (CA).</p> <p>(72) Inventors: SCHACHTER, Harry ; 5 Menin Road, Toronto, Ontario M6C 3J1 (CA). SARKAR, Mohan ; 77 Elm Street, Apartment #405, Toronto, Ontario M5G 1H4 (CA).</p> <p>(74) Agent: D'IORIO, Hélène; Gowling, Strathy &amp; Henderson, Suite 2600, 160 Elgin Street, Ottawa, Ontario K1N 8S3 (CA).</p>		<p>(81) Designated States: AT, AT (European patent), AU, BB, BE (European patent), BF (OAPI patent), BG, BJ (OAPI patent), BR, CA, CF (OAPI patent), CG (OAPI patent), CH, CH (European patent), CI (OAPI patent), CM (OAPI patent), CS, DE, DE (European patent), DK, DK (European patent), ES, ES (European patent), FI, FR (European patent), GA (OAPI patent), GB, GB (European patent), GN (OAPI patent), GR (European patent), HU, IT (European patent), JP, KP, KR, LK, LU, LU (European patent), MC, MG, ML (OAPI patent), MR (OAPI patent), MW, NL, NL (European patent), NO, PL, RO, SD, SE, SE (European patent), SN (OAPI patent), SU*, TD (OAPI patent), TG (OAPI patent).</p> <p><b>Published</b> <i>With international search report.</i></p> <p>(88) Date of publication of the international search report: 10 October 1996 (10.10.96)</p>	

(54) Title: CLONING OF UDP-N-ACETYLGLUCOSAMINE: $\alpha$ -D-MANNOSIDE  $\beta$ -1,2-N-ACETYLGLUCOSAMINYL-TRANSFERASE I



(57) Abstract

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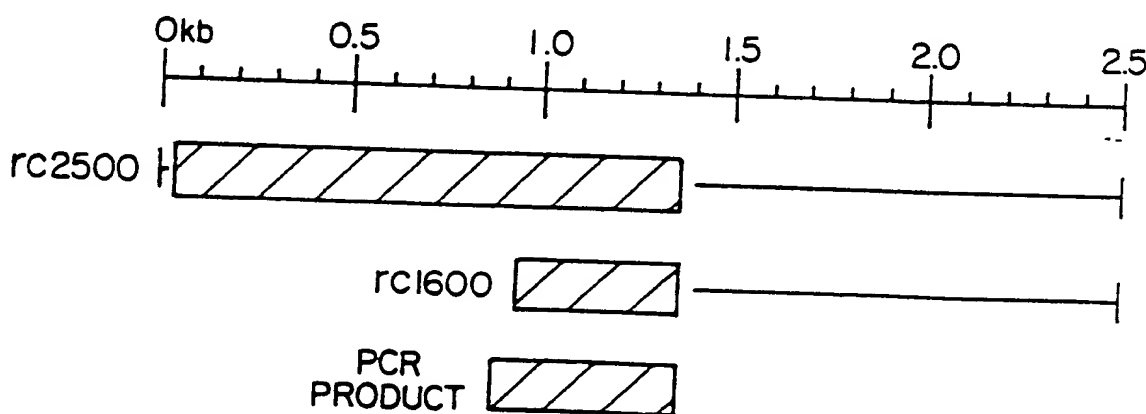


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BR	Brazil	HU	Hungary	PL	Poland
CA	Canada	IT	Italy	RO	Romania
CF	Central African Republic	JP	Japan	SD	Sudan
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CLONING OF UDP-N-ACETYLGLUCOSAMINE: $\alpha$ -3-D-MANNOSIDE  
 $\beta$ -1,2-N-ACETYLGLUCOSAMINYLTRANSFERASE I

BACKGROUND OF THE INVENTION

Field of the Invention

The present invention relates to DNA sequences for the human and rabbit enzymes which control the conversion of high mannose to hybrid and complex N-glycans, UDP-N-acetylglucosamine: $\alpha$ -3-D-mannoside  $\beta$ -1,2-N-acetylglucosaminyltransferase I (GnT I), plasmids containing such DNA sequences, transformed cells containing such plasmids, and a method for converting high mannose glycoproteins to branched N-glycan glycoproteins.

Discussion of the Background

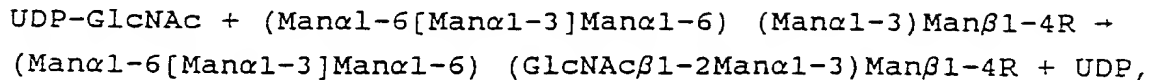
The biosynthesis of highly branched N- and O-glycans is important to many biological phenomena (Rademacher et al (1988) Ann. Rev. Biochem., vol. 57, 785-838). For example, baby hamster kidney cells transformed either by polyoma virus or by Rous sarcoma virus show a two-fold increase in one of the N-acetylglucosaminyltransferases (GlcNAc-transferase V) involved in the synthesis of highly branched complex N-glycans (Pierce et al (1986) J. Biol. Chem., vol. 261, 10772-10777; Yamashita et al (1985) J. Biol. Chem., vol. 260, 3963-3969). All N-glycans share the common core structure Man $\alpha$ 1-6(Man $\alpha$ 1-3)Man $\beta$ 1-4GlcNAc $\beta$ 1-4GlcNAc $\beta$ -Asn. Complex N-glycans have "antennae" or branches attached to this core. The antennae are initiated by the action of at least five Golgi-localized membrane-bound GlcNAc-transferases designated GnT I, II, IV, V and VI (Schachter et al (1989) Methods Enzymol., vol. 179, 351-396) and may be further elongated by the addition of D-galactose, L-fucose and sialic acid residues. Complex N-glycans may be "bisected" by a GlcNAc residue attached in  $\beta$ 1-4 linkage to the  $\beta$ -linked Man of the core due to the action of GlcNAc-transferase III (GnT III).

The conversion of high-mannose to complex and hybrid N-glycans is controlled by UDP-GlcNAc: $\alpha$ -3-D-mannoside  $\beta$ -1,2-N-

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acetylglucosaminyltransferase I (GnT I, EC 2.4.1.101), which catalyzes the reaction:



where R is  $\text{GlcNAc}\beta 1-4(+/-\text{Fuc}\alpha 1-6)\text{GlcNAc-Asn-X}$ , and Asn-X may be an Asn residue which is part of the amino acid sequence of a protein.

The enzyme is specific for the  $\text{Man}\alpha 1-3\text{Man}\beta 1-4\text{GlcNAc-arm}$  of the core. The presence of a  $\beta 2$ -linked GlcNAc residue at the non-reducing terminus of this arm is essential for subsequent action of several enzymes in the processing pathway (Schachter et al (1983) Can. J. Biochem. Cell Biol., vol. 61, 1049-1066; Schachter et al (1985)

"Glycosyltransferases involved in the biosynthesis of protein-bound oligosaccharides of the asparagine-N-acetyl-D-glucosamine and serine(threonine)-N-acetyl-D-galactosamine types", in: A.N.

Martonosi, ed. The Enzymes of Biological Membranes, New York, N.Y., Plenum Press, 227-277; Schachter, (1986)

Biochem. Cell Biol., vol. 64, 163-181; Schachter (1988)

Biochemie., vol. 70(11), 1701-1702), i.e., GnT II, III and

IV require the prior action of GnT I, and GnT V and VI require the prior action of GnT II. GnT I has been reported in hen oviduct, Chinese hamster ovary cells, baby hamster kidney cells, bovine colostrum, pig trachea and mammalian liver (Schachter et al (1983) Can. J. Biochem. Cell Biol., vol. 61, 1049-1066; Schachter et al (1985)

"Glycosyltransferases involved in the biosynthesis of protein-bound oligosaccharides of the

asparagine-N-acetyl-D-glucosamine and serine(threonine)-N-acetyl-D-galactosamine types", in: A.N. Martonosi, ed. The Enzymes of Biological Membranes, New York, N.Y., Plenum Press, 227-277; Schachter et al (1980) "Mammalian

glycosyltransferases: their role in the synthesis and function of complex carbohydrates and glycolipids", in:

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Lennarz W.J., ed. Biochemistry of Glycoproteins and Proteoglycans, New York, N.Y., Plenum Press, 85-160; Brockhausen et al (1988) Biochem. Cell Biol., vol. 66, 1134-1151). The enzyme has been partially purified from bovine colostrum (Harpaz et al (1980) J. Biol. Chem., vol. 255, 4885-4893) and from pig liver and trachea (Oppenheimer et al (1981) J. Biol. Chem., vol. 256, 11477-11482), and to homogeneity from rabbit liver (Oppenheimer et al (1981) J. Biol. Chem., vol. 256, 799-804; Nishikawa et al (1988) J. Biol. Chem., vol. 263, 8270-8281).

Recently, the cloning of DNA encoding proteins and the expression of such cloned DNA to produce the proteins has become commercially important. For ease of culturing, it is preferred that the cloned DNA be expressed in a primitive host, such as a bacteria (e.g., E. coli), a yeast, or a fungus. However, such primitive hosts may not normally possess the enzymes required for the post-translation modification of proteins which occurs in the cells from which the DNA originated. Thus, although many primitive hosts possess the necessary enzymes to effect the post-translation modification of a protein to a high mannose derivative, such host do not contain the enzyme required to convert the high mannose derivative to a hybrid and branched glycan, GnT I.

As discussed in Bergh et al, "Glycosylation of Heterologously Expressed Proteins: Problems and Solutions", in Therapeutic Peptide and Proteins: Assessing the New Technologies, Marshak et al eds, Cold Spring Harbor Laboratory, Banbury Report 29, 1988, in prokaryotes, the resulting lack of glycosylation may have a variety of consequences, such as incorrect polypeptide chain-folding, precipitation and aggregation of the protein, proteolytic degradation or enhanced immunogenicity.

Yeast and vertebrate cells use the same Glc<sub>3</sub>Man<sub>2</sub>GlcNAc<sub>2</sub> lipid-linked precursor for cotranslational glycosylation of asparagine residues, both recognize the same Asn-X-ser/Thr sequences, and both remove the three glucose residues soon

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after transfer. Thus, a mammalian glycoprotein expressed in yeast may contain the same carbohydrate chains as the native protein until after it leaves the endoplasmic reticulum. After entry into the Golgi, however, the later steps in oligosaccharide processing are very different in yeast (see Kukuruzinska et al, Ann. Rev. Biochem., vol. 56, p.915, 1987) and vertebrates, (see Hubbard and Ivatt Ann. Rev. Biochem., vol. 50, p.555, 1981; Kornfeld and Kornfeld Ann. Rev. Biochem., vol. 54, p.631, 1985). Processed Saccharomyces cerevisiae N-linked oligosaccharides contain two GlcNAc residues and from 9 to 50 or more mannose residues. On the other hand, mammalian oligosaccharides never have more than nine mannose residues and most commonly contain GlcNAc, galactose, and sialic acid attached to a Man,GlcNAc<sub>2</sub> core.

Thus, heterologous expression in yeast of a mammalian glycoprotein intended for therapeutic use can present a number of potential glycosylation-related problems. For example, carbohydrate chains may be highly antigenic; in addition, they are recognized by Man/GlcNAc-specific receptors on cells of the mammalian reticuloendothelial system, resulting in rapid clearance of the glycoprotein from the circulation.

Thus, it is desirable to: (1) provide large amounts of GnT I for the further post translational modification of recombinantly produced proteins; and (2) provide a means for enabling primitive hosts to express GnT I.

However, as yet there are no methods available for obtaining large quantities of GnT I or enabling primitive hosts to express GnT I.

#### SUMMARY OF THE INVENTION

Accordingly, it is an object of the present invention to provide a method for producing large quantities of GnT I.

It is another object to provide a method for converting high mannose derivatives to hybrid and complex N-glycans.

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It is another object to provide isolated DNA sequences which encode GnT I.

It is another object to provide plasmids which contain a DNA sequence which encodes GnT I.

It is another object to provide microorganisms which contain a heterologous sequence of DNA which encodes GnT I.

These and other objects, which will become apparent during the following detailed description, have been achieved by the inventors' isolation and cloning of DNA sequences encoding rabbit and human GnT I, preparation of plasmids containing such DNA sequences and transfection of microorganisms, with such plasmids.

#### BRIEF DESCRIPTION OF THE DRAWINGS

A more complete appreciation of the invention and many of the attendant advantages thereof will be readily obtained as the same become better understood by reference to the following detailed description when considered in connection with the accompanying drawings, wherein:

Figure 1 illustrates the amino acid sequence data for the eight peptides isolated from rabbit liver GnT I and nucleotide sequences of the six synthetic oligonucleotides prepared on the basis of the peptide sequences. The single letter code is used for amino acid sequence data; upper case letters indicate firm assignments and lower case letters indicate tentative assignments. The underlined sections of the peptide sequences indicate the regions used for the design of oligonucleotide probes. Probes 2, 3 and 6 were based on peptides 2, 3 and 6, respectively; S indicates "sense" and A indicates "antisense" directions;

Figure 2 illustrates a schematic representation of GnT I clones. PCR product, product obtained by PCR amplification of rabbit liver cDNA; rc 1600, 1.6 kb GnT I cDNA clone; rc2500, 3.0 kb GnT I cDNA clone. The shaded boxes represent the coding region. During subcloning, the 3.0 kb cDNA was reduced to 2.5 kb by a 0.5 kb deletion at the 5'-end;

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Figure 3 illustrates the results of an agarose gel electrophoresis (1% agarose) of the products of the polymerase chain reaction (PCR) using rabbit liver cDNA as template and the following combinations of oligonucleotides as primers: 2S-3A; 2S-6A; 3S-2A; 3S-6A; 6S-2A; 6S-3A (Figure 1). Conditions of PCR are given in the Methods section. The gel was stained with ethidium bromide (0.5 µg/ml). Primer-dependent products were obtained with combinations 2S-6A (0.50 kb) and 3S-6A (0.45 kb). The arrow designates the 0.5 kb DNA marker; the remaining standards are at 1.0 kb, 1.6 kb, 2.0 kb and at 1.0 kb intervals thereafter;

Figure 4 illustrates the nucleotide sequence (lower case) of the 2.5 kb GnT I cDNA clone. The amino acid sequence in the coding region is shown in upper case letters. The positions of the eight peptide sequences obtained from proteolytic digests of GnT I (Figure 1) are underlined with a single solid line; the regions of these peptide sequences used for oligonucleotide probe synthesis (Figure 1) are additionally underlined with a discontinuous line. The putative transmembrane segment (bases 62-136) is underlined with a double line. The consensus polyadenylation signal AATAAA at position 2435 is underlined. Only the nucleotide sequence is numbered;

Figure 5 illustrates an autoradiogram of an SDS-polyacrylamide gel electrophoresis experiment showing in vitro transcription and translation of the rabbit cDNA. mRNA was generated from the 2.5 kb GnT I cDNA and was used as the template for in vitro translation using rabbit reticulocyte lysate and L-[<sup>35</sup>S]-methionine (see Methods for details). Lane C, no plasmid in the incubation; lane 12, pGEM-7z containing the 2.5 kb GnT I cDNA with an insert between bases 56 and 57 which interrupts the reading frame; lane 16, pGEM-7z containing the 2.5 kb GnT I cDNA (pGEM-7z-rcgnt1);

Figure 6 illustrates the nucleotide sequence for human genomic DNA encoding for GnT I;

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Figure 7 illustrates the amino acid sequence for human GnT I; and

Figure 8 illustrates both the nucleotide sequence for human genomic DNA encoding for GnT I and the amino acid sequence of human GnT I.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Thus, one aspect of the present invention relates to isolated DNA sequences which encode rabbit GnT I. Specifically, such DNA sequences encode a protein having the sequence (starting from the N-terminal) of formula I shown below:

MET LEU LYS LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE  
LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR  
ARG PRO VAL PRO SER ARG LEU PRO SER ASP ASN ALA LEU ASP ASP  
ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP  
ALA GLU VAL GLU LEU GLU ARG GLN ARG GLY LEU LEU GLN GLN ILE  
ARG GLU HIS HIS ALA LEU TRP SER GLN ARG TRP LYS VAL PRO THR  
ALA ALA PRO PRO ALA GLN PRO HIS VAL PRO VAL THR PRO PRO PRO  
ALA VAL ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL  
ARG ARG CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU  
LEU PHE PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR  
ALA GLN VAL ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG  
GLN PRO ASP LEU SER ASN ILE ALA VAL GLN PRO ASP HIS ARG LYS  
PHE GLN GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU  
GLY GLN ILE PHE HIS ASN PHE ASN TYR PRO ALA ALA VAL VAL VAL  
GLU ASP ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE GLN  
ALA THR TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL  
SER ALA TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP SER SER  
LYS PRO GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY  
TRP LEU LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP  
PRO LYS ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG  
LYS GLY ARG ALA CYS VAL ARG PRO GLU ILE SER ARG THR MET THR  
PHE GLY ARG LYS GLY VAL SER HIS GLY GLN PHE PHE ASP GLN HIS  
LEU LYS PHE ILE LYS LEU ASN GLN GLN PHE VAL PRO PHE THR GLN  
LEU ASP LEU SER TYR LEU GLN GLN GLU ALA TYR ASP ARG ASP PHE  
LEU ALA ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL

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ARG THR ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR  
THR GLY ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL  
MET ASP ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY  
ILE VAL THR PHE LEU PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO  
PRO GLN THR TRP ASP GLY TYR ASP PRO SER TRP THR

In another aspect, the present invention relates to DNA sequences which encode human GnT I. Such DNA sequences encode a protein having the sequence (starting from the N-terminus) of formula II shown below:

1: MET LEU LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE  
16: LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR  
31: ARG PRO ALA PRO GLY ARG PRO PRO SER VAL SER ALA LEU ASP GLY  
46: ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP  
61: ALA GLU VAL GLU LEU GLU ARG ARG GLY LEU LEU GLN GLN ILE  
76: GLY ASP ALA LEU SER SER GLN ARG GLY ARG VAL PRO THR ALA ALA  
91: PRO PRO ALA GLN PRO ARG VAL PRO VAL THR PRO ALA PRO ALA VAL  
106: ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL ARG ARG  
121: CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU LEU PHE  
136: PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR ALA GLN  
151: ALA ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG GLN PRO  
166: ASP LEU SER SER ILE ALA VAL PRO PRO ASP HIS ARG LYS PHE GLN  
181: GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU GLY GLN  
196: VAL PHE ARG GLN PHE ARG PHE PRO ALA ALA VAL VAL VAL GLU ASP  
211: ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE ARG ALA THR  
226: TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL SER ALA  
241: TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP ALA SER ARG PRO  
256: GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY TRP LEU  
271: LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP PRO LYS  
286: ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG GLN GLY  
301: ARG ALA CYS ILE ARG PRO GLU ILE SER ARG THR MET THR PHE GLY  
316: ARG LYS GLY VAL THR HIS GLY GLN PHE PHE ASP GLN HIS LEU LYS  
331: PHE ILE LYS LEU ASN GLN GLN PHE VAL HIS PHE THR GLN LEU ASP  
346: LEU SER TYR LEU GLN ARG GLU ALA TYR ASP ARG ASP PHE LEU ALA  
361: ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL ARG THR  
376: ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR THR GLY  
391: ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL MET ASP  
406: ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY ILE VAL

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421: THR PHE GLN PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO PRO PRO  
436: THR TRP GLU GLY TYR ASP PRO SER TRP ASN

Exemplary of the DNA sequences encoding rabbit GnT I is  
the sequence (starting from the 5'-terminus) of formula III,  
shown below:

atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca  
cgt cca gtg cct agc agg ctg ccg tca gac aat gct ctc gat gat  
gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
gcc gag gta gag ttg gaa cgt cag ccg gga ctg ttg cag cag att  
agg gag cac cat gct ctt tgg agc cag ccg tgg aag gtg cct act  
gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
gct gtg atc ccc atc ctg gta att gcc tgt gac cgc agc acc gtc  
cgc cgc tgt ttg gac aag cta ctg cat tat cgg cct tca gct gag  
ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
gcc cag gtc att gct tcc tat ggc agc gca gtc aca cac atc ccg  
caa cct gac ctg agc aac att gct gtg cag ccc gac cac cgc aag  
ttc cag ggc tac tac aag atc gca ccg cat tac cgc tgg gca ttg  
ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg gtg  
gag gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
gcc act tac cca ctg ttg aaa gca gac ccc tcc ctc tgg tgt gtg  
tct gcc tgg aat gac aat ggc aaa gaa cag atg gta gac tcg agt  
aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggc tta ggc  
tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag ccc aag tgg  
ccc aaa gcc ttc tgg gat gac tgg atg cgc ccg cct gag cag cga  
aag ggg agg gcc tgt gtg cgt cca gaa atc tca aga aca atg aca  
ttt ggc ccg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
ctc aag ttc atc aag ctg aac cag cag ttt gta ccc ttc acc cag  
ctg gac ctg tcg tac ctt cag cag gag gcc tat gac ccg gat ttc  
ctt gct cgt gtt tat ggt gct ccc cag tta cag gtg gag aaa gtg  
agg acc aat gac ccg aag gag cta gga gag gtg cgc gta cag tac  
aca ggc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc  
atg gat gac ctc aaa tca ggt gta ccc agg gct gga tac ccg ggc  
att gtc acc ttc tta ttc ccg ggc cgc cgt gtc cac ctg ccg ccc  
cct cag act tgg gat ggc tat gat cct agt tgg act

The DNA sequence of formula III corresponds to the  
coding region of rabbit cDNA encoding GnT I. Another

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example of a DNA sequence encoding rabbit GnT I is a larger section of cDNA encoding rabbit GnT I, which has the formula IV as shown below:

1 gaattccggc aagtcatacc tttgcctgcc ctccccgtg gggggccagg  
atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca  
cgt cca gtg cct agc agg ctg ccg tca gac aat gct ctc gat gat  
gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
gcc gag gta gag ttg gaa cgt cag cggtt gga ctg ttg cag cag att  
agg gag cac cat gct ctt tgg agc cag cggtt ggg aag gtg cct act  
gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
gct gtg atc ccc atc ctg gta att gcc tgt gac cgc agc acc gtc  
cgc cgc tgt ttg gac aag cta ctg cat tat cggtt cct tca gct gag  
ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
gcc cag gtc att gct tcc tat ggcc agc gca gtc aca cac atc cggtt  
caa cct gac ctg agc aac att gct gtg cag ccg gac cac cgc aag  
ttc cag ggc tac tac aag atc gca ccg cat tac cgc tgg gca ttg  
ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg gtg  
gaa gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
gcc act tac cca ctg ttg aaa gca gac cc tcc ctc tgg tgt gtg  
tct gcc tgg aat gac aat ggcc aaa gaa cag atg gta gac tcg agt  
aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggcc tta ggc  
tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag cc aag tgg  
ccc aaa gcc ttc tgg gat gac tgg atg cgc ccg cct gag cag cga  
aag ggg agg gcc tgt gtg cgt cca gaa atc tca aga aca atg aca  
ttt ggc ccg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
ctc aag ttc atc aag ctg aac cag cag ttt gta cc ttc acc cag  
ctg gac ctg tcg tac ctt cag cag gag gcc tat gac ccg gat ttc  
ctt gct cgt gtt tat ggt gct cc cag tta cag gtg gag aaa gtg  
agg acc aat gac ccg aag gag cta gga gag gtg cgc gta cag tac  
aca ggcc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc  
atg gat gac ctc aaa tca ggt gta cc agg gct gga tac ccg ggc  
att gtc acc ttc tta ttc ccg ggc ccg cgt gtc cac ctg ccg ccc  
cct cag act tgg gat ggc tat gat cct agt tgg act  
taacagctcc tgcctgtccc ttctgggctc ctcccttgca atttcatgtat ctaagatggg  
accgttagtcc ctgggctgca ttgtcttttc tggctttccc tcttgggtcc atttttttt  
tttttttttt tgagtggcat ttgaatacac agatgacaag gtgagggttc ttttgtaaa  
ggaggttagat cagggaaagc attctgtgt ctgttgggtca tcaaggcagca aaccactgtg  
tgcatagggga agaatggct ttgtgggccc agaaatatcc atgttctgag ttttttttt

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aggtcatctg cagaggagtt ggcaacttaa gctttcttaa ccaggccttt tctttctgac  
 ctgagagcca gggcatgaga cttcttgttc atgtccctt ttaccttccc ctaataaggg  
 tctggctac aggagaagt aacatattgt ggccagaata atactaacca gaggggcctc  
 attgtcagag tctaggtgca gttattgggt tgcagagtt aatgccttct gttcttctt  
 ctttattcct gacttctgctc agcttcttct tcttgcagc ctagcaattt ttggttctaa  
 gatgaaaaat gaagaggaaa agaaatattc gcacccagct attggagaa aggtatggg  
 aaaaaaactt cattgtacca cttcaaagag acactcttga ccttccctt tctaaaaatt  
 agtccccctcc ctgttgcttc aggagaatgc tgcgtggc agttctgtgt gatecttctt  
 ccctgagttt tatacacagg ctcctcccta aggctgtggc ttctgggtgc cctctgaca  
 taagttacag tggccaagac caggacaact cccggccatga gctaagtctt gcctaccc  
 tccaaaacat tcccatgtcc tcacaggcta ggatgcagat gttgggttggg gagaaattt  
 tgtgtgtgtg tgtgtgtgtg tgtgtttct tgcctgaccc cagtttcatg gatgaaaagt  
 ggaagctaca gaattatttt caaaaataaa ggctgaattt tctgaaaaaa aaaaaaaaaa  
 aaaaaaccgg aattt

The DNA sequence of formulae III and IV have been obtained by cloning the rabbit cDNA encoding GnT I, by the procedure which is described in detail in the Examples section.

Exemplary of the DNA sequences encoding human GnT I is the sequence (starting at the 5'-terminus) of formula V, shown below:

atgctgaa gaagcagtct gcagggctt tgctgtgggg cgctatccct tttgtggcct  
 961 ggaatgcctt gctgctccctt ttcttcttggc cgcgcctcagc acctggcagg ccaccctcag  
 1021 tcagcgctct cgtatggcgtac cccggccagcc tcacccggga agtgtatcgcc ctggcccaag  
 1081 acgcccgggtt ggagctggag cgcaggcgtg ggctgtcgca gcagatcggtt gatggccctgt  
 1141 cgagccagcg ggggggggtt cccaccggcgg cccctcccgcc ccagccgggt gtgcctgtga  
 1201 ccccccggcc ggccgggtattt cccatccctgg tcatecgctg tgaccgcage actgttcggc  
 1261 gctgccttggc caagctgtcg cattatcgcc cctcggtgtt gctttttttttt atcatecgta  
 1321 gccaggactg cgggcacgag gagacggccc aggccatcgcc ctcctacggc agcgcggta  
 1381 cgcacatccg gcagcccgac ctgagcagca ttgcgggttgc gccggaccac cgcaagttcc  
 1441 agggctacta caagatcgcc cgccacttacc gctggggcgctt gggccaggc ttccggcagt  
 1501 ttcgtttttcc cgcggccgtt gttggggaggatgaccccttggg ggtggccccc gacttcttgc  
 1561 agtactttcg ggccacccat cccgtgtgttggc aggccgaccc ctcctgtgg tgcgtctcgg  
 1621 cctggaatga caacggcaag gageagatgg tggacgcacgg cagggcttggatgttcc  
 1681 gcaccgactt tttccctggc ctgggtgttgc tgctgtttggc cgagctctgg gctgagctgg  
 1741 agcccaagtg gccaaaggcc ttcttggacactggatgcg ggcggccggag cagcggcagg  
 1801 ggcggccctg catacgccctt gagatcttcaa gaacgttgcac ctttggccgc aagggtgtga  
 1861 cgcacgggca gtttttttgc cagcacctca agtttatcaa gctgaaccag cagtttgtc  
 1921 acttcacccca gctggacccat ttttacccatgc agcggggaggc ctatgaccga gatttcttgc  
 1981 ccccgctca cgggtgtttcc cagctgcagg tggagaaagt gaggaccat gaccggaaagg  
 2041 agctgggggaa ggtgcgggttgc cagttacgg ggagggaccc cttcaaggat ttcgccaagg  
 2101 ctctgggtgt tatggatgac cttaaatgcgg ggggttccgg agctggctac cggggatttgc  
 2161 tcaccccttcca gttccggggc cgcgtgttcc acctggccggcc cccaccggacg tgggagggttgc  
 2221 atgatccctatgttggat

The DNA sequence of formula V corresponds to the coding region of human genomic DNA encoding GnT I. Another example of a DNA sequence encoding human GnT I is a larger section

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of human genomic DNA encoding GnT I, which has the formula VI, shown below:

I, shown below:

1 aagttttgaa tggtaatgtt tatttaagtt tatttctaaa tttttctca tttctctggc  
61 ttttgaagt agggtttct catccatgtt ttcttcatt gagttatgg tggatatgaa  
121 ggctatccat tagtatatgt tgatttttat attacacttc cttgcgtcagt tcattattga  
181 ttcttttga gtttccagg catattctca caagttaaaga taatagaaat agtttgc  
241 ctttccactt ctgtttgaa ttttttttc ttggttcatt tgcatggct gcttcctcca  
301 gaaaaatgtt aaataaccct ggagatgtg ggcaacttcg ttttgcct gacattcgtg  
361 ggggcctct ggtgctccc tgggtttag gggtaactg tagccctgag gtgggacatt  
421 tgatttaaa aatcagtcat ctggggcgc ttaggtttaga ggaatggtag gcagatgctg  
481 tcactccctg cccctccccct cttccctccc acctggaggg gaaatgaaat ctgacaggta  
541 gaaagagggg agttggggtt cttttctct cttccctccac cagcatcaact ctgcctct  
601 ccctcaaaaa tacgttctcg gtcaggata tatgttact ccctagagag ctctggagtc  
661 aacccctgg cttccctcca ccctcactt tggcccttgc ctgccttccat ttcccttacc  
721 tggggcat ggagccacga gccttgtgt gacgggttgc ttctctcct ctgtcttttag  
781 gtcatggct gcctctaatt cccatgtcc agaggaggca tccctaggac tgcgggcaag  
841 ggagccgcaa gcccaggca gccttgaacc gtccctggc ctgcctcccg gtgggggcca  
901 ggatgctgaa gaagcagtct gcagggcttg tgctgtggg cgctatccct tttgtggc  
961 ggaatgcctt gctgttctc ttcttctgga cgcccccac acctggcagg ccaccctcag  
1021 tcagcgctct cgatggcgcac cccggcagcc tcaccggga agtgattcgc ctggcccaag  
1081 acggcgaggg ggagctggag cgcaaggcgtg ggctgtcga gcagatcggg gatccctgt  
1141 cgagccagcg ggggagggtg cccaccgcgg cccctccgc ccagcccgct gtgcctgtga  
1201 ccccccgcgc ggcgggtatt cccatctgg tcatecgctg tgaccgcagg actgttccgc  
1261 gtcgcctgga caagctgtcg cattatcggc ctcggctga gctctccccc atcatcgta  
1321 gccaggactg cgggacacgag gagacggccc aggccatcgc ctccatcggc agcggggc  
1381 cgcacatccg gcagcccgac ctgagcagca ttgcggtgcc gccggaccac cgcaagttcc  
1441 agggctacta caagatcgcg cgccactacc gtcggcgct gggccaggc ttccggc  
1501 ttcgcctccc cgccggcgtg tgggtggagg atgacctgga ggtggccccc gacttctcg  
1561 agtactttcg gcccacccat cccgtgtcga aggccgaccc ctccctgtgg tgcgtctcg  
1621 cctggatga caacggcaag gaggatgtt gggacgccc cagggctcgat ctgtcttacc  
1681 gcaccgacctt ttccctggc ctgggttggc tgctgttggc cgagctctgg gctgagctgg  
1741 agcccaagtg gccaaaggcc ttctggacg actggatcgc gggccggag cagccggcagg  
1801 ggcgggcctg catacgccct gagatctcaa gaacgatgac ctttggccgc aagggtgtga  
1861 cgcacgggca gttcttgcac cagcacccatc agtttatcaa gctgaaccag cagtttgc  
1921 acttcacccca gtcggacctg tcttacccatc acggggaggc ctatgaccga gatttctcg  
1981 cccgcgtcta cgggtgtccc cagctgcagg tggagaaagt gaggaccaat gaccggaaagg  
2041 agctggggga ggtgcgggtg cgtatacgg ggagggacag cttaaggct ttcacaggct  
2101 ctctgggtgt tatggatgac cttaaatcg tgggttccgg agctggctac cggggatttgc  
2161 tcacccatcca gttccggggc cggcgtgtcc acctggcgc cccaccgacg tgggaggc  
2221 atgatccctag ctggaaattag cacctgcctg tcccttctgg gccccttcc gccacatcat  
2281 gagctgaggt gaccacagtc cccaggcgtc atccgcctgc ctgtgttcc ctcttaggt  
2341 catttatctt ttgatttt ccgagttggca tttaaatgtc caaatgataa caagaggatt  
2401 attctccctgt ttcacaggga gtcagatcag gggactatt ctaggatgt ttgcccc  
2461 ttaagcagga aaacactgtg tgggtgggggg cactgggtt gttggggcca caaatgtcca  
2521 cgtcctgagc ttcttcctgg agcatgtcga gagatgttgg caacgttcgc tctcttgacc  
2581 agaccccttc tccctgactg gtccttccag ccaggcactga gccccttcc tatacctgc  
2641 cccctccca gtggggactg agttatggga gaaggggaca tattttgttgc caaaatgata  
2701 ctaaccaaag gggcttccctt gtcaggccct ggtggagttt gttgggttgc cttcttc  
2761 gcttcctgccc tttcttcctt gtctgacccccc cacttagccc ttccttc ttc  
2821 agtttatagt tctgagatgg aaagttgaag ggggcaagca agaccccttc tca  
2881 cccagctgtc aggagagagg tgcagggagg aaggccctt gctgggacaa cttcttc  
2941 gcttcacccctt cagagaggac tatccctga ccccttc tctgaaaatca gtc  
3001 tgggtgttgc ggggttgc gtccttgc ttttttttgc aattcgatct gcttc  
3061 ttttccctg ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3121 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3181 ttttccctg ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3241 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3301 ttttccctg ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3361 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3421 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3481 ttttccctg ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3541 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3601 ttttccctg ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3661 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3721 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3781 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3841 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3901 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
3961 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4021 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4081 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4141 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4201 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4261 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4321 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4381 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4441 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4501 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4561 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4621 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4681 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4741 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4801 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4861 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4921 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
4981 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5041 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5101 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5161 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5221 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5281 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5341 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5401 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5461 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5521 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5581 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5641 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5701 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5761 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5821 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5881 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
5941 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
6001 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc  
6061 ggggttgcaca ccccttc ttcagatcag aggtggagca gtc

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3121 ggagcagtga ccaggacgccc tctggccca tgctgcccag cctcccccgc cgctcccaagg  
3181 cgeccccatgt cctcacaggc caggacgcca tggcggccgg gaggcatgcga

The DNA sequences of formulae V and VI have been obtained by cloning human genomic DNA encoding GnT I, by the procedure which is described in detail in the Examples section.

Of course, it is to be understood that the present DNA sequences also include those which may not exactly match the sequences of formulae III-VI, but rather contain a small number of nucleotide substitutions, deletions, and/or additions. Further, the present DNA sequences also include those which encode for amino acid sequences which may not exactly match the sequences of formulae I and II, but rather contain a small number of amino acid residue substitutions, deletions, and/or additions, provided that the protein encoded by the DNA sequence exhibits GnT I activity.

In another embodiment, the present invention relates to plasmids which contain a DNA sequence encoding rabbit or human GnT I. Such plasmids may be prepared by conventional techniques and include plasmids formed by inserting one of the present DNA sequences into any suitable plasmid. Specific examples of the present plasmids include pGEM-7z-rcgnt1, in which a 2.5 kb sequence of rabbit cDNA encoding for GnT I (Figure 2) has been inserted into pGEM-7z; pGEX-2t-rcgnt1, in which a 2.5 kb sequence of rabbit cDNA encoding GnT I has been inserted into pGEX-2t; and pGEM-5z-hggnt1, in which a 4 kb sequence of human genomic DNA encoding GnT I has been inserted into pGEM-5z. The preparation of the plasmids pGEM-7z-rcgnt1, pGEX-2t-rcgnt1, and pGEM-5z-hggnt1 is described in detail in the Examples section, and all three of these plasmids have been deposited under the provisions of the Budapest Treaty with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD 20852, USA on November 30, 1990 (Accession numbers not yet known).

In another embodiment, the present invention relates to transformed microorganisms which contain a heterologous

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sequence of DNA encoding rabbit or human GnT I. Examples of suitable host cells including: bacteria, such as E. coli, Brevibacteria, and Coryneforms; fungus, such as Trichoderma reesei, Aspergillus niger, and Aspergillus awamori; yeast, such as Saccharomyces cerevisiae, Candida albicans, Candida utilis, Candida parapsilosis, Schizosaccharomyces pombe, Bandeiraea simplicifolia, Kluyveromyces lactis, Saccharomyces kluyveri, Hansenula, Saccharomyces and Pichia; and vertebrate cells such as Chinese hamster ovary cells and COS cells. The transformed cells may be prepared by transfecting the cells with any of the present plasmids by conventional methods.

Another aspect of the present invention relates to methods for the production of GnT I. In a first embodiment, the present method comprises cell-free or in vitro expression of one of the present DNA sequences to obtain GnT I. For example, in vitro transcription and translation of one of the present plasmids using a system such as described in Methods in Molecular Biology, Nucleic Acids, Walker, ed., Humana Press, Clifton, NJ, pp 145-155 (1984) yields GnT I.

In another embodiment, the present method comprises culturing a microorganism which contains a heterologous DNA sequence which corresponds to one of the present DNA sequences. Although the culturing conditions, such as time, medium, temperature, light, and agitation, will depend on the identity of the host microorganism and the yield of GnT I desired, these conditions are readily determined by those skilled in the art.

In a further aspect, the present invention relates to a method for converting a glycoprotein which is in the high mannose form to a glycoprotein which is in the form of a hybrid or complex N-glycan. In a first embodiment, the present method may be carried out by reacting, in vitro, a glycoprotein which is in the high mannose form with mannosidases followed by UDP-GlcNAc in the presence of GnT I.

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In another embodiment, the present method may comprise culturing a cell which produces a glycoprotein in high mannose form and which also contains a heterologous sequence of DNA encoding human or rabbit GnT I. For example, transfection of cell, which normally produces a glycoprotein in a mannose form, with one of the present plasmids may be used to form a cell which produces the protein (produced in high mannose form before transfection) as a hybrid or complex N-glycan. Preferably, the glycoprotein, which is produced in the high mannose form prior to transfection with the present DNA, is also produced by the host cell as a result of transformation. In other words, the DNA encoding the glycoprotein is also heterologous with respect to the host cell.

Examples of such glycoproteins are described in Tanner et al, Biochimica et Biophysica Acta; vol. 906, pp. 81-99 (1987); and Kukurazinska et al, Ann. Rev. Biochem., vol. 56, pp 915-944 (1987) and include SUC 2, CSF, c-IgM  $\mu$ -chain, c-IgM chain, c-amylase, c-HBsAg, c-hemagglutinin, c-a<sub>1</sub> antitrypsin, c-prea<sub>1</sub>, antitrypsin, c-glycoamylase, c-VSV gp, c-sindbis virus E1 gp, c-sindbis virus E2 gp, c-killerprotoxin (type I), c-phascolin  $\alpha$  and  $\beta$ , hepatitis B virus surface antigen, interferon-gamma, tissue plasminogen activator, monoclonal anti-bodies, chicken ovalbumin-like proteins, interleukin-2, and proteins from vesicular stomatitis, influenza, and Semliki Forest viruses.

As noted above, branched glycans on membrane glycoproteins have been implicated in a variety of biological phenomena, e.g. tumor progression and metastasis, embryogenesis, cell differentiation, cell-cell and receptor-ligand interactions, viral and bacterial infectivity, fertilization and the control of the immune system (Rademacher et al (1988) Ann. Rev. Biochem., vol. 57, 785-838; Pierce et al (1986) J. Biol. Chem., vol. 261, 10772-10777; Yamashita et al (1985) J. Biol. Chem., vol. 260, 3963-3969; Schachter (1986) Biochem. Cell Biol., vol. 64, 163-181; West (1986) Mol. Cell. Biochem., vol. 72,

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3-20; Narasimhan et al (1988) J. Biol. Chem., vol. 263, 1273-1281; Dennis et al (1987) Science, vol. 236, 582-585). GnT I catalyzes an essential first step in the conversion of high mannose to branched hybrid and complex N-glycans (Schachter (1986) Biochem. Cell Biol., vol. 64, 163-181; Brockhausen et al (1988) Biochem. Cell Biol., vol. 66, 1134-1151). In vitro transcription/translation of the 2.5 kb cDNA reported in this paper results in GnT I activity demonstrating the cloning of the gene for the catalytic domain of this important control enzyme.

At least seven glycosyltransferases involved in the synthesis of N- and O-glycans have been cloned to date, i.e., UDP-Gal:GlcNAc-R  $\beta$ 1,4-Gal-transferase (Appert et al (1986) Biochem. Biophys. Res. Commons., vol. 139, 163-168; D'Agostaro et al (1989) Eur. J. Biochem., vol. 183, 211-217; Masri et al (1988) Biochem. Biophys. Res. Commun., vol. 157, 657-663; Narimatsu et al (1986) Proc. Natl. Acad. Sci. USA, vol. 83, 4720-4724; Shaper et al (1986) Proc. Natl. Acad. Sci. USA, vol. 83, 1573-1577; Shaper et al (1988) J. Biol. Chem., vol. 263, 10420-10428; Nakazawa et al (1988) J. Biochem. (Tokyo), vol. 104, 165-168). UDP-Gal:Gal-R  $\alpha$ 1,3-Gal-transferase (Joziasse et al (1989) J. Biol. Chem., vol. 264, 14290-14297; Larsen et al (1989) Proc. Natl. Acad. Sci. USA, vol. 86, 8227-8231; Larsen et al (1990) J. Biol. Chem., vol. 265, 7055-7061; Smith et al (1990) J. Biol. Chem., vol. 265, 6225-6234), CMP-sialic acid:Gal-R  $\alpha$ 2,6-sialyltransferase (Weinstein et al (1987) J. Biol. Chem., vol. 262, 17735-17743), CMP-sialic acid:Gal-R  $\alpha$ 2,3-sialyltransferase (Paulson et al (1990) FASEB J., vol. 4, A1862), GDP-Fuc:Gal $\beta$ 1,4(3)GlcNAc-R (Fuc to GlcNAc) $\alpha$ 1,3(4)-Fuc-transferase (Gersten et al (1990) FASEB J., vol. 4, A1930; Kukowska-Latallo (1990) FASEB J., vol. 4, A1930), GDP-Fuc:Gal-R  $\alpha$ 1,2-Fuc-transferase (Rajan et al (1989) J. Biol. Chem., vol. 264(19), 11158-11167; Ernst et al (1989) J. Biol. Chem., vol. 264(6), 3436-3447) and UDP-GalNAc:Fuc $\alpha$ 1,2Gal-R (GalNAc to Gal)  $\alpha$ 1,3-GalNAc-transferase (Yamamoto et al (1990) J. Biol. Chem.

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Chem., vol. 265, 1146-1151). These transferases all place sugars in terminal or subterminal positions; three of them ( $\beta$ 1,4-Gal-,  $\alpha$ 2,6-sialyl-, and  $\alpha$ 1,3-GalNAc-transferases) have been localized to the trans-Golgi cisternae and trans-Golgi network, at least in some tissues (Roth et al (1982) J. Cell Biol., vol. 92, 223-229; Roth (1984) J. Cell Biol., vol. 98, 399-406; Roth (1987) Biochem. Biophys. Acta., vol. 906, 405-436; Roth et al (1988) Eur. J. Cell Biol., vol. 46, 105-112; Duncan et al (1988) J. Cell Biol., vol. 106, 617-628; Lee et al (1989) J. Biol. Chem., vol. 264, 13848-13855; Tooze et al (1988) J. Cell Biol., vol. 106, 1475-1487; Berger et al (1985) Proc. Nat. Acad. Sci. USA, vol. 82, 4736-4739; Taatjes et al (1988) J. Biol. Chem., vol. 263, 6302-6309). Human  $\alpha$ 1,3-GalNAc-transferase and a human pseudogene showing homology to murine  $\alpha$ 1,3-Gal-transferase share 55% homology (Laresen et al (1990) J. Biol. Chem., vol. 265, 7055-7061). CMP-sialic acid:Gal-R  $\alpha$ 2,6- and  $\alpha$ 2,3-sialyltransferases exhibit 50% identity and 80% conservation over a 50 amino acid stretch (Paulson et al (1990) FASEB J., vol. 4, A1862). The remaining transferases share no significant sequence similarities but have very similar domain structures, i.e., a short amino-terminal cytoplasmic tail, a 16-20 amino acid transmembrane segment (non-cleavable signal-anchor domain), a "stem" or "neck" region of undetermined length, and a long carboxyterminal catalytic domain which is in the Golgi lumen (Paulson et al (1989) J. Biol. Chem., vol. 264, 17615-17618).

The presence of a "neck" region is based on the finding that the  $\alpha$ 2,6-sialyltransferase (Weinstein et al (1987) J. Biol. Chem., vol. 262, 17735-17743; Lammers et al (1988) Biochem. J., vol. 256, 623-631) and the  $\beta$ 1,4-Gal-transferase (D'Agostaro et al (1989) Eur. J. Biochem., vol. 183, 211-217) can be cut by proteases to release a smaller catalytically active protein lacking the trans-membrane domain. The exact length of this "neck" region cannot be stated with accuracy since it is not known how much of the amino-terminal sequence can be removed without loss of

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catalytic activity. It has been shown that rabbit liver GnT I (Nishikawa et al (1988) J. Biol. Chem., vol. 263, 8270-8281) and rat liver UDP-GlcNAc: $\alpha$ -6-D-mannoside  $\beta$ -1,2-N-acetylglucosaminyltransferase II (GnT II) (Bendiak et al (1987) J. Biol. Chem., vol. 262, 5784-5790; Bendiak et al (1987) J. Biol. Chem., vol. 262, 5775-5783) exist in two forms, a large amount of presumably membrane-bound material which does not adhere to columns and a small amount of material which can be purified. In the case of GnT I, it is now clear from the sequence analysis that the 45 kDa form of the catalytically active protein previously purified has been derived from the membrane-bound precursor by proteolytic cleavage at about base position 215 in the "neck" region (Figure 4). The N-terminal blockage of this 45 kDa protein must therefore be due to chemical modification during GnT I purification. The hydrophobic trans-membrane region can form an  $\alpha$ -helix with a hydrophobic surface capable of interacting with the membrane or with other hydrophobic proteins within the membrane. This strong hydrophobic interaction may explain why it is so difficult to purify glycosyltransferase preparations with intact trans-membrane domains.

Rabbit GnT I, human, mouse and bovine UDP-Gal:GlcNAc-R  $\beta$ 1,4-Gal-transferases and human UDP-GalNAc:Fuc $\alpha$ 1,2Gal-R (GalNAc to Gal)  $\alpha$ 1,3-GalNAc-transferase have an abnormally high number of Pro residues between the transmembrane domain and the catalytic domain, e.g., there are 13 Pro residues in GnT I between the transmembrane domain and base position 376 (Figure 4); 9 of these Pro residues occur in a short stretch of 21 amino acids (bases 314-376, Figure 4). This Pro-rich "neck" may play a role in positioning the catalytic domain in the lumen of the Golgi to enable glycosylation of glycoproteins moving along the Golgi lumen.

The domain structure of GnT I appears to be similar to that of the previously cloned glycosyltransferases. However, GnT I differs from these transferases in being a medial-Golgi enzyme, at least in some tissues (Dunphy et al

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(1985) Cell, vol. 40, 463-472; Kornfeld et al (1985) Ann. Rev. Biochem., vol. 54, 631-664). Although no medial-Golgi glycosyltransferase has been cloned to date, rat liver  $\alpha$ -mannosidase II (also a medial-Golgi enzyme) has been partially cloned (Moremen (1989) Proc. Natl. Acad. Sci. USA, vol. 86(14), 5276-5280). Comparison with GnT I reveals a 16-amino acid sequence in GnT I (LHYRPSAELFPIIVSQ, bases 431-478, Figure 4) which shows a high similarity score to amino acid residues 403-418 in  $\alpha$ -mannosidase II (LQYRNQEQLFSYMNSQ). Paulson's group (Paulson et al (1989) J. Biol. Chem., vol. 264, 17615-17618; Colley et al (1989) J. Biol. Chem., vol. 264, 17619-17622) has suggested that the trans-Golgi retention signal lies in the amino-terminal 57 amino acids of the  $\alpha$ 2,6-sialyltransferase molecule. The 16-amino acid "consensus" sequence present in GnT I and  $\alpha$ -mannosidase II may be the equivalent medial-Golgi retention signal. Joziasse et al (1989) J. Biol. Chem., vol. 264, 14290-14297, have suggested that a column hexapeptide sequence K(R)DKKND(E) may serve as a UDP-Gal binding site in the  $\beta$ 1,4-Gal- and  $\alpha$ 1,3-Gal-transferases; this sequence is not present in GnT I.

Sequence data indicate that the carboxy-terminal half of human GnT I shows 87% nucleotide sequence similarity and 90% amino acid sequence similarity to the carboxy-terminal half of rabbit liver GnT I. Strong homology between species has also been observed for bovine, murine and human UDP-Gal:GlcNAc-R  $\beta$ 1,4-Gal-transferase (Appert et al (1986) Biochem. Biophys. Res. Commun., vol. 139, 163-168; D'Agostaro et al (1989) Eur. J. Biochem., vol 183, 211-217; Masri et al (1988) Biochem. Biophys. Res. Commun., vol. 157, 657-663; Narimatsu et al (1986) Proc. Nat. Acad. Sci. USA, vol. 83, 4720-4724; Shaper et al (1986) Proc. Nat. Acad. Sci. USA, vol. 83, 1573-1577; Shaper et al (1988) J. Biol. Chem., vol. 263, 10420-10428; Nakazawa et al (1988) J. Biochem. (Tokyo), vol. 104, 165-168) bovine and murine UDP-Gal:Gal-R  $\alpha$ 1,3-Gal-transferase (Joziasse et al (1989) J. Biol. Chem., vol. 264, 14290-14297; Larsen et al (1989)

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Proc. Natl. Acad. Sci. USA, vol. 86, 8227-8231), murine and human GDP-Fuc:Gal $\beta$ 1,4(3)GlcNAc-R (Fuc to GlcNAc)  $\alpha$ 1,3(4)-Fuc-transferase (Gersten et al (1990) FASEB J., vol. 4, A1930; Kukowska-Latallo et al (1990) FASEB J., vol. 4, A1930), and human and rat CMP-sialic acid:Gal-R  $\alpha$ 2,6-sialyltransferase (Lance et al (1989) Biochem. Biophys. Res. Commun., vol. 164, 225-232).

It has been reported (Kumar et al (1990) Mol. Cell Biol., vol. 9, 5713-5717; Ripka et al (1989) Biochem. Biophys. Res. Commun., vol. 159(2), 554-560; Ripka et al (1990) J. Cellular Biochem., vol. 42, 117-122) that transformation of Lec I Chinese hamster ovary (CHO) cell mutants (which lack GnT I) with a crude preparation of total human genomic DNA results in transfectants expressing GnT I enzyme activity; this approach should allow cloning of the human GnT I gene by the gene transfer and expression screening method recently used to clone several glycosyltransferases (Larsen et al (1989) Proc. Natl. Acad. Sci. USA, vol. 86, 8227-8231; Larsen et al (1990) J. Biol. Chem., vol. 265, 7055-7061; Smith et al (1990) J. Biol. Chem., vol. 265, 6225-6234; Gersten (1990) FASEB J., vol. 4, A1930; Kukowska-Latallo et al (1990) FASEB J., vol. 4, A1930; Rajan et al (1989) J. Biol. Chem., vol. 264(19), 11158-11167; Ernst et al (1989) J. Biol. Chem., vol. 264(6), 3436-3447).

Other features of the invention will become apparent in the course of the following descriptions of exemplary embodiments which are given for illustration of the invention and are not intended to be limiting thereof.

#### EXAMPLES

##### I. Rabbit:

Preparation of Peptides. Rabbit liver GnT I was purified as previously described (Nishikawa et al (1988) J. Biol. Chem., vol. 263, 8270-8281). Glycerol, Triton X-100 and salts were removed from the purified enzyme (approximately 15  $\mu$ g) by "inverse-gradient" reversed-phase

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high performance liquid chromatography (RP-HPLC) (Simpson et al (1987) Eur. J. Biochem., vol. 165, 21-29). The enzyme solution (100  $\mu$ l) was diluted to 1.2 ml with n-propanol in a sample-loading syringe, thoroughly mixed, and loaded at 1 ml/min on a VeloSep C<sub>8</sub> cartridge (3- $\mu$ m particle size, 30 x 2.1 mm i.d.; Applied Biosystems, Foster City, CA, USA) previously equilibrated in 100% n-propanol at 40°C. GnT I was retained on the reversed-phase column under these conditions whereas glycerol, Triton X-100 and salts were washed through the column with 100% n-propanol. GnT I was eluted at 0.1 ml/min as a sharp peak by a linear gradient (5%/min) of decreasing n-propanol concentration (100% to 50%) generated with 100% n-propanol and 50% n-propanol/50% water containing 0.4% (v/v) trifluoroacetic acid at 40°C. GnT I-containing fractions from the inverse gradient RP-HPLC were pooled, adjusted to 0.02% (w/v) with respect to Tween 20 (Pierce Chemical Co., Rockford, IL, USA), concentrated to 100  $\mu$ l in a 1.5-ml polypropylene tube using a centrifugal vacuum concentrator to reduce the n-propanol concentration, and diluted to 1.5 ml with 5% (v/v) formic acid containing 0.02% Tween 20.

Edman degradation of purified GnT I ( $\sim$  200 pmol) yielded no N-terminal sequence indicating N-terminal blockage; proteolysis of GnT I was therefore undertaken. GnT I was digested with pepsin (Sigma) at an enzyme/substrate mass ratio of 1:20 for 1 h at 37°C and the digest was fractionated by RP-HPLC on a short microbore column (30 x 2.1 mm i.d.) employing a low pH (trifluoroacetic acid, pH 2.1) mobile phase and a gradient of acetonitrile to yield peptides 5 and 6 (Figure 1). Core GnT I remaining after pepsin digestion was reduced with dithiothreitol and alkylated with iodoacetic acid (Simpson et al (1988) Eur. J. Biochem., vol. 176, 187-197) to give core S-carboxymethylated(SCM)-GnT I which was purified by RP-HPLC (Simpson et al (1988) Eur. J. Biochem., vol. 176, 187-197; Simpson et al (1989) Anal. Biochem., vol. 177, 221-236). Pepsin-treated core SCM-GnT I (about 10  $\mu$ g in

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1 ml 1% ammonium bicarbonate, 1mM CaCl<sub>2</sub>, 0.02% Tween 20) was digested with trypsin (Worthington) at an enzyme/substrate mass ratio of 1:20 for 16 h at 37°C. RP-HPLC of the digest showed that trypsin resulted in little further digestion of the pepsin-treated material. Sequence analysis of a portion of this material resulted in 33 amino acid assignments (peptide 1, Figure 1). Pepsin and trypsin-treated core SCM-GnT I (about 8 µg in 1 ml 1% ammonium bicarbonate-0.02% Tween 20) was digested with thermolysin (Sigma) at an enzyme/substrate mass ratio of 1:20 for 2 h at 50°C and the digest was fractionated by RP-HPLC to yield peptides 2, 3, 4, 7 and 8 (Figure 1). Core GnT I was extremely resistant to proteolysis even after reduction and alkylation indicating that the molecule is probably very compact.

HPLC. RP-HPLC was carried out on a Hewlett-Packard liquid chromatograph (model 1090A) fitted with a diode array detector (model 1040A) (Simpson et al (1988) Eur. J. Biochem., vol. 176, 187-197). A Brownlee RP-300 column (30-nm pore size, 7-µm diameter dimethyloctylsilica particles packed into a stainless steel cartridge, 30 x 2.1 mm i.d.; Brownlee Laboratories, Santa Clara, CA, USA) was used for all peptide separations.

Amino Acid Sequence Analysis. Automated amino acid sequence analysis of GnT I and derived peptides was performed with Applied Biosystems sequencers (models 470A and 477A) equipped with on-line phenylthiohydantoin (PTH) amino acid analyzers (model 120A). Polybrene (Klapper et al (1978) Anal. Biochem., vol. 85, 126-131) was used as a carrier.

Oligonucleotides and cDNA Synthesis. Oligonucleotides were synthesized on a Pharmacia automated oligonucleotide synthesizer at the Hospital for Sick Children-Pharmacia Biotechnology Service Centre. Total RNA was prepared from rabbit liver by the method of Chirgwin et al (Chirgwin et al (1979) Biochemistry, vol. 18, 5294-5299; Ausubel et al (1990) Current Protocols in Molecular Biology, Media, PA:Greene Publishing Associates and John Wiley and Sons).

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Poly(A)+RNA was prepared by oligo(dt) chromatography (Aviv et al (1972) Proc. Natl. Acad. Sci. USA, vol. 69, 1408-1412) using the mRNA Purification Kit supplied by Pharmacia. Single-stranded cDNA synthesis was performed using the RiboClone cDNA Synthesis System (Promega) with the following modifications. Total rabbit liver RNA (20 µg) in a volume of 5.5 µl was heated at 65°C for 3 min followed by cooling on ice for 5 min. The following reagents were added to a final volume of 50 µl: 50 mM Tris-HCl, pH 8.3; 0.15 M KCl; 10 mM MgCl<sub>2</sub>; 2 mM dithiothreitol (DTT); each dNTP at 0.4 mM; 40 units of RNasin (Promega); 2 mM sodium pyrophosphate; a mixture of the three anti-sense oligonucleotide primers 2A, 3A and 6A (Figure 1) at concentrations of 50 nM each; 20 units of AMV reverse transcriptase and 15 units of murine leukemia virus reverse transcriptase. Incubation was at 42°C for 2 hr. The reaction mixture was treated with NaOH (0.25 N final concentration) for 5 min at room temperature to destroy RNA. The solution was then heated at 65°C for 1 min followed by cooling on ice for 5 min and neutralized with HCl (0.25 N final concentration). This cDNA preparation was used directly in the PCR reaction.

Amplification of cDNA. PCR was carried out in a total volume of 0.1 ml containing 50 mM KCl, 10 mM Tris-HCl (pH 8.3), 1.5 mM MgCl<sub>2</sub>, 0.01% gelatin, each of the four dNTP at 0.2 mM, 0.5 µM of each oligonucleotide in six paired combinations of oligonucleotide primers (2S-3A, 2S-6A, 3S-2A, 3S-6A, 6S-2A, 6S-3A, Figure 1), 10 µl of RNA-free rabbit liver cDNA (see above), 2.5 units of *Thermus aquaticus* (Taq) polymerase (Perkin-Elmer/Cetus) and 0.1 ml of mineral oil. The samples were placed in an automated heating/cooling block (DNA Thermal Cycler, Perkin-Elmer) programmed for a temperature-step cycle of 94°C (0.5 min), 50°C (1 min) and 72°C (2 min) for a total of 40 cycles followed by a 10-minute extension at 72°C after the final cycle. DNA from the PCR reactions was purified with GeneClean (Bio 101, Inc.) and analyzed by electrophoresis in a 1% agarose gel containing ethidium bromide (0.5 µg/ml).

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Two PCR products (0.45 and 0.50 kb) were detected and were purified from a 1% agarose gel by GeneClean. The DNA ends were filled in with T4 DNA polymerase (Moremen (1989) Proc. Natl. Acad. Sci. USA, vol. 86(14), 5276-5280) and the blunt ends were ligated into SmaI site of pGEM-7z (Promega). The recombinant plasmid was amplified in E. coli XL1-blue cells and purified. The plasmid was used for sequencing and to prepare a labelled probe for screening of a cDNA library.

Screening of rabbit liver cDNA library in λgt10. The recombinant plasmid containing pGEM-7z and 0.5 kb PCR product (see above) was cut with BamH1 and used to generate a riboprobe (0.5 kb) with the Promega Riboprobe Gemini II Core System. The reaction contained in a total volume of 25 μl:32 mM Tris-HCl, pH 7.5; 5 mM MgCl<sub>2</sub>; 2 mM spermidine; 8 mM sodium chloride; 8 mM DTT; 40 units RNasin; 0.4 mM of each of ATP, GTP and UTP; 5 μl[α-<sup>32</sup>P]CTP (800 Ci/mmol); 1 μg of BamH1-cut pGEM-7z/PCR-product recombinant plasmid; and 2 units T7 RNA polymerase. Incubation was at 40°C for 2 hr. RNase-free DNase I (10 units) was added followed by incubation at room temperature for 15 min. Buffer (80 μl of 50 mM Tris-HCl, pH 7.4; 4 mM EDTA; 300 mM NaCl; 0.1% SDS) and tRNA (20 μg) were added followed by extraction with phenol-chloroform-isoamyl alcohol (25:24:1, v/v). The labelled RNA probe was desalting over a Sephadex G-50 column (Nick Column, Pharmacia).

A rabbit liver cDNA library in λgt 10 (5'-stretch, Cat. No TL 1006a from Clontech, EcoRI cloning site) was propagated in E. coli LE392 host cells and 10<sup>6</sup> plaques were screened by standard plaque hybridization techniques (Maniatis et al (1982) Molecular Cloning: a laboratory manual, Cold Spring Harbor, N.Y.:Cold Spring Harbor Laboratory) using the above riboprobe. Following fixation of DNA to nitrocellulose membranes, the membranes were washed for 1 hr at 45°C in 50 mM Tris-HCl, pH 8.0/1 M NaCl/1 mM EDTA/0.1% SDS. Membranes were prehybridized at 50°C for 2 hr in 1M NaCl/50 mM sodium phosphate, pH 6.5/0.1% SDS/50% freshly-deionized formamide/1% glycine/0.5% Blotto/5 mM

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EDTA/1% yeast total RNA. Riboprobe ( $5 \times 10^6$  cpm/ml hybridization solution) was added and hybridization was carried out at 50°C overnight. Membranes were washed in 2XSSC/0.1% SDS twice for 5 min at room temperature and twice for 15 min at 50°C. Positive isolates were identified by autoradiography and were plaque-purified. DNA was purified from phage lysates, digested with EcoRI, and cDNA inserts were analyzed by agarose gel electrophoresis. The largest cDNA insert obtained was 1.6 kb; it was subcloned into the EcoRI site of pGEM-7z (Promega) by standard methods (Maniatis et al (1982) Molecular Cloning: a laboratory manual, Cold Spring Harbor, N.Y.:Cold Spring Harbor Laboratory) and the recombinant plasmids were transfected into E. coli XL1-blue. Colonies containing the recombinant plasmid were selected and amplified, and plasmid DNA was purified by CsCl gradient centrifugation (Ausubel et al (1990) Current Protocols in Molecular Biology, Media, PA:Greene Publishing Associates and John Wiley and Sons).

The cDNA library was re-screened as described above using a 80 bp riboprobe prepared from the 5'-end of the 1.6 kb clone. The largest cDNA insert obtained was 3.0 kb. This insert was sub-cloned into pGEM-7z as described above and plasmid DNA was purified by CsCl gradient centrifugation (Ausubel et al (1990) Current Protocols in Molecular Biology. Media, PA:Greene Publishing Associates and John Wiley and Sons), to obtain pGEM-7z-rcgnt1.

DNA Sequencing. Two colonies of the pGEM-7z/PCR-product recombinant plasmid (see above) containing inserts in opposite directions were sequenced directly by the single-strand dideoxynucleotide-chain-termination method (Sanger et al, Proc. Natl. Acad. Sci. USA, vol. 74, 5463-5467) using deoxyadenosine 5'-[ $\alpha$ -[<sup>35</sup>S]thio] triphosphate, Sequenase (United States Biochemical) and the forward primer for pGEM-7z. The 1.6 and 3.0 kb clones were sequenced by the Erase-a-Base System (Promega) and the single-strand dideoxynucleotide-chain-termination method. Both DNA strands were sequenced by using colonies in which

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the inserts were present in opposite directions. Plasmid DNA (12 µg) was cut with SphI to generate a 5'-overhang and XbaI to generate a 3'-overhang. The cut DNA was digested with exonuclease III (Erase-a-Base System, Promega) for varying lengths of time followed by S1 nuclease digestion. The DNA ends were blunt-ended with the Klenow fragment of E. coli DNA polymerase I and the DNA was circularized with T4 DNA ligase. The ligation mixtures were transfected into competent XL1-blue cells. Miniplasmid preparations were carried out on about 5-10 subclones from each exonuclease III time point and were cut with BamHI and AatII to determine DNA size. Colonies with appropriate deletions were amplified and incubated with M13KO7 helper phage at 37°C for 1 hr followed by amplification in the presence of kanamycin (70 µg/ml) for 6 hr at 37°C. Single-stranded DNA was produced by the helper phage and excreted into the medium. The ss-DNA was purified from the medium by polyethylene glycol precipitation and sequenced by the dideoxynucleotide chain-termination method using deoxyadenosine 5'-[ $\alpha$ -[<sup>35</sup>S]thio]triphosphate, Sequenase (United States Biochemical) and the forward primer for pGEM-7z.

RNA Hybridization. Rabbit liver poly(A)+RNA (5 µg) was denatured in 50% (v/v) formamide/6% (v/v) formaldehyde buffer at 65°C and was resolved by gel electrophoresis in a 1% agarose gel containing 6% (v/v) formaldehyde. The RNA was transferred to a nitrocellulose filter and the filters were hybridized with the <sup>32</sup>P-labelled 0.5 kb PCR riboprobe (see above) followed by autoradiography. The specific activity of the probe was about 10<sup>6</sup> dpm/ng and the hybridization solution contained about 10<sup>6</sup> dpm/ml.

In vitro transcription and translation. The recombinant plasmid containing pGEM-7z (Promega) and the 2.5 kb Gnt I cDNA insert (rc2500, Figure 2) (pGEM-7z-rcgnt1) was cut with Sph I to generate linear plasmid. RNA was transcribed using the SP6 RNA polymerase promoter and initiation site present in pGEM-7z. RNA synthesis was

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carried out at 40°C for 1 hr in a total volume of 50 µl containing 40 mM Tris-HCl (pH 7.5), 6 mM MgCl<sub>2</sub>, 2 mM spermidine, 10 mM NaCl, 10 mM DTT, 40 units RNasin (Promega), 0.5 mM of each of ATP, UTP and CTP, 0.1 mM GTP, 0.5 mM m<sup>7</sup>G(5')PPP(5')G (Pharmacia), 10 units SP6 RNA polymerase and 10 µg linearized plasmid. Control incubations were carried out in the absence of plasmid or with a linearized pGEM-7z recombinant plasmid containing a non-coding insert. The reaction mixture was extracted twice with phenol-chloroform-isoamyl alcohol (25:24:1, v/v) followed by precipitation with cold ethanol.

Protein synthesis (translation) was carried out at 30°C for 1 hr in a total volume of 50 µl containing all 20 amino acids (1 mM each), 20 units of RNasin, RNA as prepared above, and buffer and rabbit reticulocyte lysate as supplied by Promega (Oliver et al (1984) "In vitro translation of messenger RNA in a rabbit reticulocyte lysate cell-free system", in: M. Walker J., ed., Methods in Molecular Biology, Nucleic Acids, Clifton, N.J.:Humana Press, 145-155). Non-radioactive amino acids were used when the products of translation were assayed for GnT I activity (see below). Separate incubations were carried out with L-[<sup>35</sup>S]-methionine (1000 Ci/mmol; 90 µCi/incubation) replacing non-radioactive Met; these incubations were analyzed by SDS-polyacrylamide gel electrophoresis followed by autoradiography.

GnT I was assayed (Schachter (1989) Methods Enzymol., vol. 179, 351-396; Brockhausen et al (1988) Biochem. Cell Biol., vol. 66, 1134-1151) in a total volume of 40 µl containing 20 mM MnCl<sub>2</sub>, bovine serum albumin (1 mg/ml), 0.1% (v/v) Triton X-100, 0.1 M MES (pH 6.1), 0.5 mM UDP-N-[1-<sup>14</sup>C]acetyl-D-glucosamine (2.2 mCi/mmol), 0.125 M GlcNAc and 0.6 mM Man $\alpha$ 1-6(Man $\alpha$ 1-3)Man $\beta$ -hexyl (a kind gift from Dr. Hans Paulsen, University of Hamburg, Hamburg, Federal Republic of Germany). Incubations were at 37°C for 2 and 16 hr. The reaction was stopped with 0.5 ml 20 mM sodium tetraborate/2 mM EDTA and was passed through a small column of AG1X8

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(Cl-form, 100-200 mesh, equilibrated with water) to remove radioactive nucleotide-sugar. The eluate was applied to a Sep-Pak C-18 reverse phase cartridge (Waters) conditioned with 20 ml methanol and 20 ml water. The cartridge was washed with 20 ml water and radioactive product was eluted with 5.0 ml methanol (Palcic et al (1988) Glycoconjugate J., vol. 5, 49-63). An aliquot was counted directly and the remainder was analyzed by HPLC on a C-18 reverse phase column using acetonitrile-water (12:88) as the mobile phase (Schachter et al (1989) Methods Enzymol., vol. 179, 351-396; Brockhausen et al (1988) Biochem. Cell Biol., vol. 66, 1134-1151). Product co-eluted with a standard preparation of Man $\alpha$ 1-6(GlcNAc $\beta$ 1-2Man $\alpha$ 1-3)Man $\beta$ -hexyl at 36 min.

Preparation of pGEX-2t-rcgnt1. This plasmid was prepared from PGEM-7z-rcgnt1 by cutting out the insert rrgnt1 with Eco RI. Plasmid pGEX-2t (Pharmacia) was linearized with Eco RI and the insert was ligated into the plasmid by standard procedures. The recombinant plasmid was amplified in E. coli in the presence of ampicillin and purified by cesium chloride centrifugation.

Amplification of cDNA. Three amino acid sequences (Figure 1) were chosen for the design of sense and anti-sense oligonucleotide primers to be used in the PCR amplification of rabbit liver cDNA. Deoxyinosine was substituted in positions where codon degeneracy was >2 (Moremen (1989) Proc. Natl. Acad. Sci. USA, vol. 85(14), 5276-5280); mixed pairs of bases were used in four positions in all three sequences giving a 16-fold mixture of sequences for every primer. Since we had no knowledge of the order of the peptides in the amino acid sequence, PCR was carried out with all six possible combinations of sense and anti-sense primers (2S-3A, 2S-6A, 3S-2A, 3S-6A, 6S-2A, 6S-3A, Figure 1). The products of the PCR reactions were analyzed by agarose gel electrophoresis (Figure 3). Primer-dependent products were obtained with two of the six incubations, i.e., 2S-6A (500 bp) and 3S-6A (450 bp). The complete nucleotide sequence for Gnt I is shown in Figure 4.

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Oligonucleotide primers 2S and 3A are separated by only nine bases thereby explaining the absence of PCR product with this combination.

Sequence Analysis. The 1.6 kb clone contains 0.5 kb from the 3'-end of the coding region and the full 1.1 kb 3'-untranslated region (rc1600, Figure 2). The 3.0 kb clone yielded a 2485 bp sequence (rc2500, Figure 2; Figure 4). We have shown that subcloning of the 3.0 kb DNA fragment in pGEM-7z results in deletion of a 0.5 kb DNA fragment near the 5'-end of the clone. Comparison of the cDNA sequence shown in Figure 4 with the sequence of human genomic DNA for GnT I (in preparation) has shown that this deleted 0.5 kb DNA fragment is not part of the GnT I gene; we do not know the origin of this DNA.

The GnT I coding sequence has 1341 bp and codes for a membrane-bound protein of 447 amino acids ( $M_r$  52,000). There is a single hydrophobic domain (bases 62 to 136) flanked by charged amino acids (Figure 4). Chou-Fasman rules (Chou et al (1978) Adv. Enzymol., vol. 47, 45-147) predict that this hydrophobic segment is capable of propagating an  $\alpha$ -helix, as expected for a transmembrane domain.

The presumptive initiation Met codon is at the ATG codon at position 50 which has an A at position 47 thereby fulfilling the requirements for an initiation codon (Kozak (1983) Microbiological Reviews, vol. 47, 1-45). All eight peptides shown in Figure 1 (a total of 103 amino acid residues) can be identified in the sequence (Figure 4); an additional five tentative assignments also match the sequence. GnT I purified from rabbit liver has a molecular weight of about 45 kDa (Nishikawa et al (1988) J. Biol. Chem., vol. 263, 8270-8281). The protein has no N-glycans since none of the nine Asn residues are in a typical Asn-X-Ser(Thr) sequence; we have previously shown that rabbit liver GnT I binds poorly to lectin/agarose columns (Nishikawa et al (1988) J. Biol. Chem., vol. 263, 8270-8281). If there are no or few O-glycans, a

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catalytically active protein of 45 kDa can be derived by cleavage at about base position 215 (Figure 4).

Comparison of the GnT I sequence with those of several previously cloned glycosyltransferases (Appert et al (1986) Biochem. Biophys. Res. Commun., vol. 139, 163-168; D'Agostaro et al (1989) Eur. J. Biochem., vol. 183, 211-217; Hollis et al (1989) Biochem. Biophys. Res. Commun., vol. 162, 1069-1075; Joziasse et al (1989) J. Biol. Chem., vol. 264, 14290-14297; Larsen et al (1989) Proc. Natl. Acad. Sci. USA, vol. 86, 8227-8231; Larsen et al (1990) J. Biol. Chem., vol. 265, 7055-7061; Masibay et al (1989) Proc. Natl. Acad. Sci. USA, vol. 86, 5733-5737; Masri et al (1988) Biochem. Biophys. Res. Commun., vol. 157, 657-663; Narimatsu et al (1986) Proc. Nat. Acad. Sci. USA, vol. 83, 4720-4724; Russo et al (1990) J. Biol. Chem., vol. 265, 3324-3331; Shaper et al (1986) Proc. Nat. Acad. Sci. USA, vol. 83, 1573-1577; Shaper et al (1988) J. Biol. Chem., vol. 263, 10420-10428; Shaper et al (1988) Biochemie., vol. 70, 1683-1688; Shaper et al (1990) Proc. Natl. Acad. Sci. USA, vol. 87, 791-795; Smith et al (1990) J. Biol. Chem., vol. 265, 6225-6234; Weinstein et al (1987) J. Biol. Chem., vol. 262, 17735-17743) revealed no sequence homology but GnT I appears to have a domain structure typical of these enzymes (Paulson et al (1989) J. Biol. Chem., vol. 264, 17615-17618). Searches of the GenBank nucleotide data base (release 62.0) with the coding region of GnT I and of the PIR Protein Data Base (release 23.0) with the GnT I amino acid sequence revealed no significant similarities to other sequences.

The complete sequence has a long 3'-untranslated region (bases 1391-2479) containing the consensus polyadenylation signal AATAAA at position 2435 (Tosi et al (1981) Nucleic Acids Research, vol. 9, 2313-2323). Long 3'-untranslated regions are typical of the known glycosyltransferase genes and may be a feature present in other Golgi-localized enzymes (Moremen (1989) Proc. Natl. Acad. Sci. USA, vol. 86(14), 5276-5280).

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Northern Blot Analysis. The PCR riboprobe was used to determine the size of mRNA in rabbit liver. A major band was detected at about 3.0 kb with some smearing at lower molecular weights (data not shown) indicating that the 2.5 kb cDNA clone (Figure 4) may not be full-length.

In Vitro transcription and translation. Transcription of the linearized pGEM-7z/2.5 kb GnT I cDNA recombinant plasmid (pGEM-7z-rcgnt1) followed by translation in the presence of L-[<sup>35</sup>S]Met resulted in the appearance of a strong radioactive 52 kDa band on SDS-polyacrylamide gel electrophoresis; this band was not seen in control incubations lacking plasmid or containing control plasmid (Figure 5). The molecular weight matches the prediction for the open reading frame shown in Figure 4. Table 1 shows the results of GnT I assays carried out on the transcription-translation incubations. The incubation containing the pGEM-7z/2.5 kb GnT I cDNA recombinant plasmid (pGEM-7z-rcgnt1) has appreciable GnT I activity whereas both controls show low activity. It is concluded that the 2.5 kb sequence shown in Figure 4 can code for the synthesis of catalytically active GnT I.

TABLE 1

In vitro transcription-translation of rabbit GnT I cDNA

Conditions of transcription	GnT I product (nmoles/total transcription incubation)		
	Sep-Pak assays 2 hr      16 hr		HPLC assays 16 hr
No plasmid	0.04	0.21	--
Control Plasmid	0.04	0.21	0.29
2.5 kb GnT I cDNA (pGEM-7z-rcgnt1)	0.41	1.05	1.32

II. Human GnT I:

The polymerase chain reaction (PCR) was used to obtain a 0.5 kb ds-cDNA representing the carboxy terminal half of the rabbit liver GnT I coding sequence and labelled this DNA fragment by the random primer technique. The preparation of this probe is described above.

The rabbit cDNA probe was used to screen  $10^6$  plaques from an amplified human genomic DNA library in  $\lambda$ EMBL3 prepared from chromosomal DNA from chronic myeloid leukemia cells. Positive plaques (23) were purified and phage DNA was subjected to restriction enzyme analysis using the 0.5 kb rabbit cDNA as probe. All 23 preparations gave the same Sau3A 0.4 kb fragment. This fragment showed 87% base similarity and 90% amino acid sequence similarity to the rabbit GnT I carboxy-terminal sequence. Inserts of 13 and 15 kb were cut from two of the human genomic DNA clones with SAI and subcloned into plasmid pGEM-5zf(+) (Promega). Restriction maps of the two inserts show that they represent an over-lapping 18 kb DNA sequence.

The coding sequence was located in a 4.0 kb fragment of human genomic DNA by screening restriction maps with a probe containing the entire coding region of the rabbit GnT I cDNA. This 4.0 kb DNA fragment was cut out by restriction enzymes and subcloned into the sequencing vector pGEM-5zf(+) to yield pGEM-5z-hggnt1 and sequenced. Transfection of the gene into Lec 1 Chinese hamster ovary cell mutants (which lack GnT I activity) results in the expression of GnT I activity indicating the presence of a functional promoter 5'-upstream of the transcription start site.

The 4 kb sequence contains an open reading frame coding for a protein with 445 amino acids (2 less than the rabbit enzyme). The DNA contains a functional promoter and an intronless gene. The similarity between the rabbit and human enzymes is 85% for the nucleotide coding sequences and over 90% for the amino acid sequences.

Obviously, numerous modifications and variations of the present invention are possible in light of the above

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teachings. It is therefore to be understood that, within the scope of the appended claims, the invention may be practiced otherwise than as specifically described herein. The references cited in the specification are incorporated herein by reference.

THE EMBODIMENTS OF THE INVENTION IN WHICH AN EXCLUSIVE PROPERTY OR PRIVILEGE IS CLAIMED ARE DEFINED AS FOLLOWS:

1. An isolated DNA sequence encoding a protein having the  
amino acid sequence of formula I:

amino acid sequence of formula 1.

MET LEU LYS LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE  
LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR  
ARG PRO VAL PRO SER ARG LEU PRO SER ASP ASN ALA LEU ASP ASP  
ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP  
ALA GLU VAL GLU LEU GLU ARG GLN ARG GLY LEU LEU GLN GLN ILE  
ARG GLU HIS HIS ALA LEU TRP SER GLN ARG TRP LYS VAL PRO THR  
ALA ALA PRO PRO ALA GLN PRO HIS VAL PRO VAL THR PRO PRO PRO  
ALA VAL ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL  
ARG ARG CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU  
LEU PHE PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR  
ALA GLN VAL ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG  
GLN PRO ASP LEU SER ASN ILE ALA VAL GLN PRO ASP HIS ARG LYS  
PHE GLN GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU  
GLY GLN ILE PHE HIS ASN PHE ASN TYR PRO ALA ALA VAL VAL VAL  
GLU ASP ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE GLN  
ALA THR TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL  
SER ALA TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP SER SER  
LYS PRO GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY  
TRP LEU LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP  
PRO LYS ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG  
LYS GLY ARG ALA CYS VAL ARG PRO GLU ILE SER ARG THR MET THR  
PHE GLY ARG LYS GLY VAL SER HIS GLY GLN PHE PHE ASP GLN HIS  
LEU LYS PHE ILE LYS LEU ASN GLN GLN PHE VAL PRO PHE THR GLN  
LEU ASP LEU SER TYR LEU GLN GLN GLU ALA TYR ASP ARG ASP PHE  
LEU ALA ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL  
ARG THR ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR  
THR GLY ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL  
MET ASP ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY  
ILE VAL THR PHE LEU PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO  
PRO GLN THR TRP ASP GLY TYR ASP PRO SER TRP THR.

2. The DNA sequence of Claim 1, having the nucleotide sequence of formula III:

sequence of formula III:

atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca

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cgt cca gtg cct agc agg ctg ccg tca gac aat gct ctc gat gat  
gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
gcc gag gta gag ttg gaa cgt cag ccg gga ctg ttg cag cag att  
agg gag cac cat gct ctt tgg agc cag ccg tgg aag gtg cct act  
gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
gct gtg atc ccc atc ctc gta att gcc tgt gac cgc agc acc gtc  
cgc cgc tgt ttg gac aag cta ctg cat tat ccg cct tca gct gag  
ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
gcc cag gtc att gct tcc tat ggc agc gca gtc aca cac atc ccg  
caa cct gac ctg agc aac att gct gtg cag ccc gac cac cgc aag  
ttc cag ggc tac tac aag atc gca ccg cat tac cgc tgg gca ttg  
ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg gtg  
gag gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
gcc act tac cca ctg ttg aaa gca gac ccc tcc ctc tgg tgt gtg  
tct gcc tgg aat gac aat ggc aaa gaa cag atg gta gac tcg agt  
aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggc tta ggc  
tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag ccc aag tgg  
ccc aaa gcc ttc tgg gat gac tgg atg cgc ccg cct gag cag cga  
aag ggg agg gcc tgg cgt cca gaa atc tca aga aca atg aca  
ttt ggc ccg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
ctc aag ttc atc aag ctg aac cag cag ttt gta ccc ttc acc cag  
ctg gac ctg tcg tac ctt cag cag gag gcc tat gac ccg gat ttc  
ctt gct cgt gtt tat ggt gct ccc cag tta cag gtg gag aaa gtg  
agg acc aat gac ccg aag gag cta gga gag gtg cgc gta cag tac  
aca ggc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc  
atg gat gac ctc aaa tca ggt gta ccc agg gct gga tac ccg ggc  
att gtc acc ttc tta ttc ccg ggc cgc cgt gtc cac ctg gcg ccc  
cct cag act tgg gat ggc tat gat cct agt tgg act.

3. The DNA sequence of Claim 1, having the nucleotide sequence of formula IV:

gaattccggc aagtcatacc tttgcctgcc ctccccctgtg gggggccagg  
atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca  
cgt cca gtg cct agc agg ctg ccg tca gac aat gct ctc gat gat  
gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
gcc gag gta gag ttg gaa cgt cag ccg gga ctg ttg cag cag att  
agg gag cac cat gct ctt tgg agc cag ccg tgg aag gtg cct act

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gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
 gct gtg atc ccc atc ctg gta att gcc tgt gac cgc agc acc gtc  
 cgc cgc tgt ttg gac aag cta ctg cat tat cgg cct tca gct gag  
 ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
 gcc cag gtc att gct tcc tat ggc agc gca gtc aca cac atc cgg  
 caa cct gac ctg agc aac att gct gtg cag ccc gac cac cgc aag  
 ttc cag ggc tac tac aag atc gca cgg cat tac cgc tgg gca ttg  
 ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg  
 gaa gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
 gcc act tac cca ctg ttg aaa gca gac ccc tcc ctc tgg tgt gtg  
 tct gcc tgg aat gac aat ggc aaa gaa cag atg gta gac tcg agt  
 aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggc tta ggc  
 tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag ccc aag tgg  
 ccc aaa gcc ttc tgg gat gac tgg atg cgc cgg cct gag cag cga  
 aag ggg agg gcc tgt gtg cgt cca gaa atc tca aga aca atg aca  
 ttt ggc cgg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
 ctc aag ttc atc aag ctg aac cag cag ttt gta ccc ttc acc cag  
 ctg gac ctg tcg tac ctt cag cag gag gcc tat gac cgg gat ttc  
 ctt gct cgt gtt tat ggt gct ccc cag tta cag gtg gag aaa gtg  
 agg acc aat gac cgg aag gag cta gga gag gtg cgc gta cag tac  
 aca ggc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc  
 atg gat gac ctc aaa tca ggt gta ccc agg gct gga tac cgg ggc  
 att gtc acc ttc tta ttc cgg ggc cgc cgt gtc cac ctg gcg ccc  
 cct cag act tgg gat ggc tat gat cct agt tgg act  
 taacagctcc tgccctgtcccc ttctgggctc ttcttgca atttcatgtat ctaagatggg  
 accgtagtcc ctgggctgca ttgtcttttc tgcctttccc tcttgggctc atttttttt  
 tttttttttt tgagtggcat ttgaatacac agatgacaag gtgagggttc ttttgtaaa  
 ggagtttagat cagggaaagc attctgctgt ctgttggta tcaagcagca aaccactgtg  
 tgatagggga agaatgggct ttttggggcc agaaatatcc atgttctgag ttttcttctt  
 aggtcatctg cagaggagtt ggcaacttta gcttcttaa ccaggcctt tcttcttgac  
 ctgagagcca gggcatgaga cttcttggtc atgetcctt ttaccttccc ctaataaggg  
 tctgggctac aggagaagtg aacatattgt ggccagaata atactaacca gagggggctc  
 attgtcagag tcttaggtgca gttattgggt tgcagatgat aatgccttctt gttttttttt  
 ccttattttt gacttctgtc agctttttt tctttgcage ctagcaattt ttggttctaa  
 gataaaaat gaagaggaaa agaaatattt gcacccagat attggggagaa aggttagtggg  
 aaaaaaaaaactt cattgtacca cttcaaagag acacttttgat ccttttttctt tctaaaaattt  
 agtccccctcc ctgggcttc aggagaatgc tgcgtggc agttctgtgt gatccttctt  
 ccctgagttt tatacacagg cttcttcccta aggctgtggc ttctgggtggc cctcctgaca  
 taagttacag tggccaagac caggacaact ccggccatga gctaagttctt gcctacctt  
 tccaaaaacat tcccatgtcc tcacaggcta ggatgcagat gttgggttggg gagaatttt  
 tgcgtgtgtg tgcgtgtgtg tgcgtttttt tgcctgacat cagtttcatg gatgaaaaatgg  
 ggaagctaca gaattttttt caaaaaataaa ggctgaattt tctgaaaaaaaaaaaaaaa  
 aaaaaaaaaaccgg aatcc.

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4. An isolated DNA sequence encoding a protein having the amino acid sequence of formula II:

MET LEU LYS LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR ARG PRO ALA PRO GLY ARG PRO PRO SER VAL SER ALA LEU ASP GLY ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP ALA GLU VAL GLU LEU GLU ARG ARG GLY LEU LEU GLN GLN ILE GLY ASP ALA LEU SER SER GLN ARG GLY ARG VAL PRO THR ALA ALA PRO PRO ALA GLN PRO ARG VAL PRO VAL THR PRO ALA PRO ALA VAL ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL ARG ARG CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU LEU PHE PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR ALA GLN ALA ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG GLN PRO ASP LEU SER SER ILE ALA VAL PRO PRO ASP HIS ARG LYS PHE GLN GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU GLY GLN VAL PHE ARG GLN PHE ARG PHE PRO ALA ALA VAL VAL GLU ASP ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE ARG ALA THR TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL SER ALA TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP ALA SER ARG PRO GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY TRP LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP PRO LYS ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG GLN GLY ARG ALA CYS ILE ARG PRO GLU ILE SER ARG THR MET THR PHE GLY ARG LYS GLY VAL THR HIS GLY GLN PHE PHE ASP GLN HIS LEU LYS PHE ILE LYS LEU ASN GLN GLN PHE VAL HIS PHE THR GLN LEU ASP LEU SER TYR LEU GLN ARG GLU ALA TYR ASP ARG ASP PHE LEU ALA ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL ARG THR ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR THR GLY ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL MET ASP ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY ILE VAL THR PHE GLN PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO PRO PRO.

5. The DNA sequence of Claim 4, having the nucleotide sequence of formula V:

atgctgaa gaaggcagtct gcagggcttg tgctgtgggg cgctatccctc tttgtggcct ggaatgcctt gctgctccct ttctttctggta cgcgcggcagc acctggcagg ccaccctcag tcagcgctct ctagggcgac cccgcagcc tcacccggga agtgattcgc ctggcccaag acggccgaggt ggagctggag cgcaggcgtg ggctgtcgca gcagatcggt gatgccctgt cgagccagcg ggggagggtg cccacccggcgg cccctccgc ccagccgcgt gtgcctgtga ccccccgcc ggcgggtgatt cccatccctgg tcatacgctg tgaccgcage actgttcggc

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gctgcctgga caagctgctg cattatcgcc cctcggctga gctttcccc atcatcgta  
 gccaggactg cgggcacgag gagacggccc aggccatcgcc ctctacggc agcgcggtca  
 cgcacatccg gcagcccgac ctgagcagca ttgcggtgcc gcccggaccac cgcaagttcc  
 aggctacta caagatcgcg cgccactacc gctggcgct ggccagggtc ttccggcagt  
 ttgcgttccc cgccggcgtg gtgggtggagg atgacctgga ggtggccccc gacttctcg  
 agtactttcg ggccacctat ccgctgctga aggccgaccc ctccctgtgg tgcgtctcg  
 cctggaatga caacggcaag gaggcagatgg tggacgccag caggcctgag ctgctctacc  
 gcaccgactt ttccctggc ctgggctggc tgctgtggc cgagctctgg gctgagctgg  
 agcccaagtg gccaaaggcc ttctgggacg actggatgcg gcccggag cagcggcagg  
 ggcggccctg catacgccct gagatctcaa gaacgatgac ctttggccgc aagggtgtga  
 cgcacggca gttcttgcg cagcaccta agtttatcaa gctgaaccag cagtttgtgc  
 acttcaccca gctggacctg ttttacctgc agcgggaggc ctatgaccga gatttctcg  
 ccccgctcta cggtgcctcc cagctgcagg tggagaaaagt gaggaccaat gaccggaaagg  
 agctggggga ggtgcgggtg cagtatacgg ggagggacag ctcaggctt tcgcacaagg  
 ctctgggtgt tatggatgac cttaaatgcgg gggttccgag agctggctac cggggatttg  
 tcacccctcca gttccggggc cgccgtgtcc acctggcgc cccaccgacg tgggagggt  
 atgatccttag ctggaaat

6. The DNA sequence of Claim 4, having the nucleotide sequence of formula VI:

aagttttgaa tgtttaagtt tatttaagtt tattttctaa tttttctca tttctctggc  
 ttttgcgtt agggttttctt catccatgtt ttctctcat gatgttatttggatattgaa  
 ggctatccat tagtatatgt tgatttttat attacacttc cttgctcgt tcattattga  
 ttcttttgcgtt gtttccagg catattctca caagtaaaga taatagaat agtttgcctc  
 ctttccaccc ttgcgttgcgaa ttttttttc ttgggttcatt tgcatggct gcttcccca  
 gcaaaatgtt aaataaccctt ggagatgatggcaacttcg ttttgcctc gacattcgtg  
 ggggtgcctt ggtgccttccc ttgggttaag gggtaactg tagccctgag gtgggacatt  
 tgattttaaa aatcaagtcat ttggggcgc tttaggtttaga ggaatggtag gcagatgctg  
 tcactccttgc cccctccccc cttcccttccc acctggaggg gaaatgaat ctgacaggta  
 gaaagagggg agttgggggtt ctttttcttctt cttccctccac cagcatcaact ctctgcctt  
 ccctcaaaaaa tacgttccctg ggtcaggata tatgttact ccctagagag ctctggagtc  
 aaccccttgc cttccctcca ccctcaactt tggcttttc ctgcctccat ttcccttacc  
 tgggggcattt gtagccacga cccttgcgtt gacggtttgc tttctcttc ctgtcttttag  
 gtgcattggcgtt ccctccataat cccatagttc agaggaggca cccttaggac tgccggcaag  
 ggagccgcaaa gcccaggcga cccttgcacc gtccctggc ctgcctcccg gtggggccca  
 ggatgctgaa gaagcagtctt cccttgcgtt tgctgtgggg cgctatcctc tttgtggccct  
 ggaatgcctt gctgccttccc ttcttcttgcg ccgcggccagg acctggcagg ccacccctcag  
 tcagcgttccctcgatggcgcac ccgcggccagg tcacccggga agtgcattgc ctggcccaag  
 acggccgaggt ggagctggag ccgcggccgtt ggctgtgcgc gcaatcggtt gatggccctgt  
 ccggccaggccgggggtt ccacccggccgg cccttccgc ccagccgcgt gtgcctgtga  
 ccccccggcc ggcgggtattt ccacatcttgc tcatcgcttgc tgaccgcagg actgttccggc  
 gctgccttgcgca caagctgcttgc cattatcgcc cctcggctga gcttttttttccatcgta  
 gcccggactt cgggcacgag gagacggccc aggccatcgcc ctccctacggc agcgcggtca  
 cgcacatccg gcagcccgac ctgagcagca ttgcggtgcc gcccggaccac cgcaagttcc  
 aggctacta caagatcgcg cgccactacc gctggcgctt gggccagggtc ttccggcagt  
 ttgcgttccc cgccggccgtt gtgggtggagg atgacctgga ggtggccccc gacttctcg  
 agtactttcg ggccacctat ccgctgctga aggccgaccc ctccctgtgg tgcgtctcg  
 cctggaatga caacggcaag gaggcagatgg tggacgccag caggcctgag ctgctctacc  
 gcaccgactt ttccctggc ctgggctggc tgctgtttggc cgagctctgg gctgagctgg  
 agcccaagtg gccaaaggcc ttctgggacg actggatgcg gcccggag cagcggcagg  
 ggccggccctg catacgccct gagatctcaa gaacgatgac ctttggccgc aagggtgtga  
 cgccacggca gttcttgcg cagcaccta agtttatcaa gctgaaccag cagtttgtgc  
 acttcaccca gctggacctg ttttacctgc agcgggaggc ctatgaccga gatttccctcg  
 ccccgctcta cggtgcctcc cagctgcagg tggagaaaagt gaggaccaat gaccggaaagg  
 agctggggga ggtgcgggtt cagtatacgg ggagggacag cttcaaggct tcgcacaagg

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ctctgggtgt tatggatgac cttaaagtccgg gggttccgag agctggctac cggggatttg  
 tcaccccca gttccggggc cgccgtgtcc acctggcgcc cccaccgcac tgggagggtt  
 atgatccctag ctggaaattag cacctgcctg tccttcctgg gccccttctt gccacatcat  
 gagctgaggt gaccacagtcccaggctgc atcggcctgc ctgtgtttcc ctcttaggtg  
 catttatctt tttgattttt ccagatggca tttaagtgc caaatgataa caagaggatt  
 attctccctt tctcaaggga gtcatcgatggactattt ctagggatgtt ttgcggggta  
 ttaaggcagga aaacactgtg tggtgggggg cactgggctt gttggggcca caaatgtcca  
 cgtcctgagc tttctcctgg agcatgtgc gagatgttgg caacgttcgc tctcttgacc  
 agaccccttc tccctgactg gctcttccag ccaggcacga gcctcccttc tataacctgct  
 ccccttccca gtggggactg agttatggga gaaggggaca tatttgcgc caaaatgata  
 ctaaccaaag gggcttcctt gtcaggccc ggtggagttt gtgggtcactc ggggctcact  
 gcctccctgccc cttcttcctt gtctgaccccc cacttagcccc ttctcttcctt gcagcctagc  
 agttatagtt tctgagatgg aaagttgaag gggggcaagca agaccttc tcaagccatg  
 cccagctgtc aggagagagg tgcaggagg aaggcctt gctgggacaa cctcttcctt  
 gccttacctt cagagaggac tatgccttga cccctccctt ctgaaaatca gtgcctccct  
 tggtgtctta ggaggcttccct gctggcttgg tagaagacag aattcgtatctt gcctgtccct  
 ttttccctgtt ggggttgaca cacaggctcc tctcagcatg aggtggagca gtgaccaggt  
 ggagcagtga ccaggacgcc tctggccctt gctggccctt cctcccccgc cgtccccagg  
 cgccccatgtt cttcacagggc caggacgcac tggcggccgg gagcatgcga.

7. A plasmid, comprising a DNA sequence encoding a protein having the amino acid sequence of formula I:

MET LEU LYS LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE  
 LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR  
 ARG PRO VAL PRO SER ARG LEU PRO SER ASP ASN ALA LEU ASP ASP  
 ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP  
 ALA GLU VAL GLU LEU GLU ARG GLN ARG GLY LEU LEU GLN GLN ILE  
 ARG GLU HIS HIS ALA LEU TRP SER GLN ARG TRP LYS VAL PRO THR  
 ALA ALA PRO PRO ALA GLN PRO HIS VAL PRO VAL THR PRO PRO PRO  
 ALA VAL ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL  
 ARG ARG CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU  
 LEU PHE PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR  
 ALA GLN VAL ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG  
 GLN PRO ASP LEU SER ASN ILE ALA VAL GLN PRO ASP HIS ARG LYS  
 PHE GLN GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU  
 GLY GLN ILE PHE HIS ASN PHE ASN TYR PRO ALA ALA VAL VAL VAL  
 GLU ASP ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE GLN  
 ALA THR TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL  
 SER ALA TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP SER SER  
 LYS PRO GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY  
 TRP LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP  
 PRO LYS ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG  
 LYS GLY ARG ALA CYS VAL ARG PRO GLU ILE SER ARG THR MET THR  
 PHE GLY ARG LYS GLY VAL SER HIS GLY GLN PHE PHE ASP GLN HIS

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LEU LYS PHE ILE LYS LEU ASN GLN GLN PHE VAL PRO PHE THR GLN  
LEU ASP LEU SER TYR LEU GLN GLN GLU ALA TYR ASP ARG ASP PHE  
LEU ALA ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL  
ARG THR ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR  
THR GLY ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL  
MET ASP ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY  
ILE VAL THR PHE LEU PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO  
PRO GLN THR TRP ASP GLY TYR ASP PRO SER TRP THR.

8. The plasmid of Claim 7, wherein said DNA sequence has  
the formula III:

atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca  
cgt cca gtg cct agc agg ctg ccg tca gac aat gct ctc gat gat  
gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
gcc gag gta gag ttg gaa cgt cag cgg gga ctg ttg cag cag att  
agg gag cac cat gct ctt tgg agc cag cgg tgg aag gtg cct act  
gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
gct gtg atc ccc atc ctg gta att gcc tgt gac cgc agc acc gtc  
cgc cgc tgt ttg gac aag cta ctg cat tat cgg cct tca gct gag  
ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
gcc cag gtc att gct tcc tat ggc agc gca gtc aca cac atc cgg  
caa cct gac ctg agc aac att gct gtg cag ccc gac cac cgc aag  
ttc cag ggc tac tac aag atc gca cgg cat tac cgc tgg gca ttg  
ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg  
gag gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
gcc act tac cca ctg ttg aaa gca gac ccc tcc ctc tgg tgt gtg  
tct gcc tgg aat gac aat ggc aaa gaa cag atg gta gac tcg agt  
aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggc tta ggc  
tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag ccc aag tgg  
ccc aaa gcc ttc tgg gat gac tgg atg cgc cgg cct gag cag cga  
aag ggg agg gcc tgt gtg cgt cca gaa atc tca aga aca atg aca  
ttt ggc cgg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
ctc aag ttc atc aag ctg aac cag cag ttt gta ccc ttc acc cag  
ctg gac ctg tcg tac ctt cag cag gag gcc tat gac cgg gat ttc  
ctt gct cgt gtt tat ggt gct ccc cag tta cag gtg gag aaa gtg  
agg acc aat gac cgg aag gag cta gga gag gtg cgc gta cag tac  
aca ggc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc

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atg gat gac ctc aaa tca ggt gta ccc agg gct gga tac cgg ggc  
att gtc acc ttc tta ttc cgg ggc cgc cgt gtc cac ctg gcg ccc  
cct cag act tgg gat ggc tat gat cct agt tgg act.

9. The plasmid of Claim 7, wherein said DNA sequence has  
the formula IV:

gaattccggc aagtcatacc tttgcctgcc ctccccctgtg gggggccagg  
atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca  
cgt cca gtg cct agc agg ctg cgg tca gac aat gct ctc gat gat  
gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
gcc gag gta gag ttg gaa cgt cag cgg gga ctg ttg cag cag att  
agg gag cac cat gct ctt tgg agc cag cgg tgg aag gtg cct act  
gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
gct gtg atc ccc atc ctg gta att gcc tgt gac cgc agc acc gtc  
cgc cgc tgt ttg gac aag cta ctg cat tat cgg cct tca gct gag  
ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
gcc cag gtc att gct tcc tat ggc agc gca gtc aca cac atc ccg  
caa cct gac ctg agc aac att gct gtg cag ccc gac cac cgc aag  
ttc cag ggc tac tac aag atc gca cgg cat tac cgc tgg gca ttg  
ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg gtg  
gaa gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
gcc act tac cca ctg ttg aaa gca gac ccc tcc ctc tgg tgt gtg  
tct gcc tgg aat gac aat ggc aaa gaa cag atg gta gac tcg agt  
aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggc tta ggc  
tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag ccc aag tgg  
ccc aaa gcc ttc tgg gat gac tgg atg cgc cgg cct gag cag cga  
aag ggg agg gcc tgt gtg cgt cca gaa atc tca aga aca atg aca  
ttt ggc cgg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
ctc aag ttc atc aag ctg aac cag cag ttt gta ccc ttc acc cag  
ctg gac ctg tcg tac ctt cag cag gag gcc tat gac cgg gat ttc  
ctt gct cgt gtt tat ggt gct ccc cag tta cag gtg gag aaa gtg  
agg acc aat gac cgg aag gag cta gga gag gtg cgc gta cag tac  
aca ggc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc  
atg gat gac ctc aaa tca ggt gta ccc agg gct gga tac cgg ggc  
att gtc acc ttc tta ttc cgg ggc cgc cgt gtc cac ctg gcg ccc  
cct cag act tgg gat ggc tat gat cct agt tgg act

taacagctcc tgcctgtccc ttctgggctc cttccttgc aatttcatgtat ctaagatggg  
 accgttagtcc ctgggctgca ttgtcttttc tgcctttcc tcttgggtcc atttttttt  
 ttttctttt tgagtggcat ttgaatacac agatgacaag gtgagggttc ttttgttaaa  
 ggaggttagat cagggaaagc attctgctgt ctgttggta tcaagcagca aaccactgtg  
 tgatagggga agaatggct ttttggggcc agaaatatcc atgttctgag ttttcttctt  
 aggtcatctg cagaggagtt ggcaacttta getttctta ccaggccttt ttttctgac  
 ctgagagcca gggcatgaga cttcttgc atgtccctt ttaccttccc ctaataaggg  
 tctggctac aggagaagtg aacatattgt ggccagaata atactaacca gaggggcctc  
 attgtcagag tcttaggtgca gttattgggt tgtcagagtt aatgccttctt gttcttctt  
 ccttatttcct gacttctgtc agtcttctt tcttgcagc ctagcaattt ttgggtctaa  
 gataaaaat gaagagggaaa agaaatattc gcacccagct attgggagaa aggttagtggg  
 aaaaaaactt cattgtacca cttcaaagag acacttcttga ccttccctt tctaaaaattt  
 agccccctcc ctgttgcctt aggagaatgc tgcgtggc agtctgtgtt gatccttctt  
 ccctgagttt tatacacagg ctctcccta aggctgtggc ttctggtgcc cctcctgaca  
 taagttacag tggccaaagac caggacaact cggccatga gctaagtccctt gcctaccc  
 tccaaaacat tcccatgtcc tcacaggcta ggatgcagat gttggttgga gaggaattt  
 tgcgtgtgtg tgcgtgtgtg tgcgtgtgtg tgcgtgtgtg tgcgtgtgtg tgcgtgtgtg  
 ggaagctaca gaattttttt caaaaataaaa ggctgaattt tctgaaaaaaaaaaaaaaa  
 aaaaaaccgg aattt.

10. A plasmid, comprising a DNA sequence encoding a protein having the amino acid sequence of formula II:

MET LEU LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE  
 LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR  
 ARG PRO ALA PRO GLY ARG PRO PRO SER VAL SER ALA LEU ASP GLY  
 ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP  
 ALA GLU VAL GLU LEU GLU ARG ARG GLY LEU LEU GLN GLN ILE  
 GLY ASP ALA LEU SER SER GLN ARG GLY ARG VAL PRO THR ALA ALA  
 PRO PRO ALA GLN PRO ARG VAL PRO VAL THR PRO ALA PRO ALA VAL  
 ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL ARG ARG  
 CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU LEU PHE  
 PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR ALA GLN  
 ALA ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG GLN PRO  
 ASP LEU SER SER ILE ALA VAL PRO PRO ASP HIS ARG LYS PHE GLN  
 GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU GLY GLN  
 VAL PHE ARG GLN PHE ARG PHE PRO ALA ALA VAL VAL GLU ASP  
 ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE ARG ALA THR  
 TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL SER ALA  
 TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP ALA SER ARG PRO  
 GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY TRP LEU  
 LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP PRO LYS  
 ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG GLN GLY  
 ARG ALA CYS ILE ARG PRO GLU ILE SER ARG THR MET THR PHE GLY  
 ARG LYS GLY VAL THR HIS GLY GLN PHE PHE ASP GLN HIS LEU LYS

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PHE ILE LYS LEU ASN GLN GLN PHE VAL HIS PHE THR GLN LEU ASP  
LEU SER TYR LEU GLN ARG GLU ALA TYR ASP ARG ASP PHE LEU ALA  
ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL ARG THR  
ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR THR GLY  
ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL MET ASP  
ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY ILE VAL  
THR PHE GLN PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO PRO PRO.  
11. The plasmid of Claim 10, wherein said DNA sequence has  
the formula V:

atgctgaa gaaggcagtct gcagggcttg tgctgtgggg cgctatccctc tttgtggcct  
ggaatgcctc gctgctccctc ttcttctggta cgcgccccagc acctggcagg ccaccctcag  
tcagegcctc cgatggcgcac cccgccagcc tcacccggga agtgattcgc ctggcccaag  
acgcccgggt ggagctggag cgcaggcgtg ggctgctgca gcagatcggg gatgccctgt  
cgagccagcg ggggagggtg cccaccgcgg cccctccgc ccagccgcgt gtgcctgtga  
ccccccgcgc ggcgggtgatt cccatcctgg tcatcgcctg tgaccgcgc actgttcggc  
gctgcctgga caagctgctg cattatcgcc cctcggctga gctttcccc atcatcgta  
gccaggactg cgggcacgag gagacggccc aggccatcgc ctcctacggc agcgggtca  
cgcacatccg gcagccgcac ctgagcagaca ttgcgggtgcc gccggaccac cgcaagttcc  
agggctacta caagatcgcg cgccactacc gctgggcgcgt gggccagggtc ttccggcagt  
ttcgcttccc cgccggcgtg gtgggtggagg atgacctggta ggtggccccc gacttctcg  
agtactttcg ggccacctat cgcgtctgaa aggccgacccc ctcctgtgg tgctctcg  
cctggaatga caacggcaag gaggcagatgg tggacgcgcag caggcctgag ctgctctacc  
gcacccgactt ttccctggc ctggggctggc tgctgttggc cgagctctgg gctgagctgg  
agcccaagtg gccaaaggcc ttctgggacg actggatgcg gccggccggag cagccggcagg  
ggccggcctg catacgccct gagatctcaa gaacgatgac ctttggccgc aagggtgtga  
cgcacggcga gttctttgac cagcacctca agtttatcaa gctgaaccag cagtttgtgc  
acttcaccca gctggacctg tcttacctgc agcgggaggc ctatgaccga gatttctcg  
cccgcgctca cggtgcgtccc cagctgcagg tggagaaaat gaggaccaat gaccggaaagg  
agctggggga ggtgcgggtg cagtatacgg ggagggacag cttcaaggct ttccccaagg  
ctctgggtgt tatggatgac cttaaatcggtt gggttcccgag agctggctac cggggatttg  
tcacccctcca gttccggggc cggcgtgtcc acctggcgc cccaccgacg tggagggct  
atgatccat ctggaa.

12. The plasmid of Claim 10, wherein said DNA sequence has  
the formula VI:

aagttttgaa tggtaatgtt tatttctaaa tattttctca tttctctggc  
ttttgttaatgtt agggtttct catccatgtt ttcttctcat gagttatgg tggatatgaa  
ggctatccat tagtataatgt tgattttat attacacttc cttgtctcgt tcattattga  
ttctttttaa gtttccagg catattctca caagtaaaga taatagaaat agtttgcctc  
cttccaccc ctgttttgc ttttttttc ttgttttcatt tgcattggct gcttccctca  
gaaaaatgtt aaataaccct ggagatgatg ggcaacttcg ttttgcctt gacatcg  
gggtgcctt ggtgcctccc tggtaatgtt gggtaactg tagccctgag gtgggacatt  
tgattttaa aatcagtcat cttggggcgc ttaggttaga ggaatggtag gcagatgc  
tcactccttgc cccctccctt cctccctccc acctggaggg gaaatgaaat ctgacaggt  
gaaagagggg agttgggggtt cttttctct ctcctccac cagcatcaact ctctgcctct  
ccctcaaaaa tacgttctg ggtcaggata tattttactt ccctagagag ctctggagtc  
aacctccctgg ccttccctca ccctcaactt tggccctttc ctgcggccat ttctcttacc  
tgtggggcat ggagccacga gcctttgtgt gacggtttgc ttctctctc ctgtcttttag  
gtgcattggct gccttcaat cccatagtc agaggaggca tccctaggac tgcggggcaag  
ggagccgcaaa gcccaggcga gccttgaacc gtcccccggc ctgcggcccg gtggggggcca  
ggatgctgaa gaaggcagtct gcagggcttg tgctgtgggg cgctatccctc tttgtggcct

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ggaatggccct gctgctcctc ttcttcgtga cgccccagc acctggcagg ccaccctca  
tcagcgctct ccatggcgac cccgcagcc tcacccggga agtgattcgc ctggcccaag  
acgcccggagt ggagctggag cgaggcggtg ggctgtcga gcagatcgaa gatggccctgt  
cgagccagcg ggggggggtg cccacccggg cccctcccg ccagccggt gtgcctgtga  
ccccccgcgc ggcgggtatt cccatctgg tcatgcctg tgaccgcagc actgttcggc  
gctgcttggaa caagctgtg cattatcgcc cctcggctga gctttcccc atcatcgta  
gccaggactg cgggcacggag gagacggccc aggccatcgc ctccatcgcc agcgcggc  
cgcacatccg ctagcccgac ctgagcagca ttgcgggtcc gcccggaccac cgcaagtccc  
agggctacta caagatcgcc cgccactacc gctggggcgt gggccaggc ttccggc  
ttcgcttccc cgccggccgt gtggggagg atgacctgaa ggtggccccg gacttttc  
agtactttcg ggcacccat cccgtgtcga aggccgaccc cccctgtgg tgctctcgg  
ccttggaaatga caacggcaag gagcagatgg tggaccccg caggcctgag ctgctctacc  
gcacccgactt ttccctggc ctgggctggc tgctgttggc caggtctgg gctgagctgg  
agcccaagtg gccaaaggcc ttctgggacg actggatgcg gcccggagg cagcggcagg  
ggcggccctg catacgccct gagatctcaa gaacgatgac ctttggccgc aagggtgtga  
cgcacccggca gttctttgac cagcacctca agtttatcaa gctgaaccag cagtttgc  
acttccaccca gctggacccg tcttacctgc agcggggaggc ctatgaccga gatttcc  
cccgcgctca cgggtgtccc cagctgcagg tggagaaaat gaggaccaat gaccggagg  
agctggggga ggtgcgggtg cagatacgg ggagggacag cttcaaggct ttcgccaagg  
ctctgggtgt tatggatgac cttaaatcgg gggttccgag agctggctac cggggatttgc  
tcacccctca gttccggggc cggcggttcc acctggcgcc cccaccgacg tgggagggt  
atgatccctag ctggaaatgg cacctgcctg tccttcgtgg gccccttctt gccacatcat  
gagctgaggc gaccacagtc cccaggctgc atcggcctgc ctgtgtttcc ctcttaggt  
catttatctt tttgattttt ccgagtggca ttaatgtca caatgataa caagaggatt  
attctccctg tctcaaggga gtcagatcag gggaaactatt ctaggatgt tgcggggta  
ttaaggcagga aaacactgtg tgggggggg cactggcctt gttggggcca caaatgtcca  
cgctctgagc tttctctgg agcatgtca gagagtttgg caacgttcgc tctcttgacc  
agaccccttc tccctgactg gctttccag ccaggcacga gccttccttc tatactgct  
ccccctccca gtggggactg agttatggga gaaggggaca tatttgtggc caaaatgata  
ctaaccaaag gggcttcctt gtcaaggccct ggtggagttg gtgggtcate ggggctcact  
gcctctgccc ctctctcttc gtctgacccc cacttagccc ttcttcctt gcagccatgc  
agtttatagt tctgagatgg aaagttgaag ggggcaagca agacccctcc teagcccatg  
cccagctgtc aggagagagg tgcaaggagg aaggccctgt gctgggacaa cctcttc  
gccttaccc ttagccctga ccccttcctt ctgaaaaatca gtgccttc  
tgtgtctca ggaggcttc gctggcttgg tagaagacag aattcgatct gcctgtcc  
tttccctg gggtttggaca cacaggctcc tctcagcatg agtggagca gtgaccagg  
ggagcagtga ccaggacgccc tctggccctag tgctgcggcag cctcccccgc cgctcc  
cgccccatgt cctcacaggc caggacgcca tggcgccgg gacatgcga.

13. A transformed cell, containing a heterologous sequence of DNA encoding a protein having the amino acid sequence of formula I.

14. The transformed cell of Claim 13, wherein said cell has the formula III.

15. The transformed cell of Claim 13, wherein said cell has the formula IV.

15. The transformed heterologous DNA sequence has the formula IV.

16. A transformed cell, containing a heterologous sequence of DNA encoding a protein having the amino acid sequence of

formula II.

... transformed cell of Claim 16, wherein said

17. The transformed cell of Claim 16, where  
heterologous DNA sequence has the formula V.

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18. The transformed cell of Claim 16, wherein said heterologous DNA sequence has the formula VI.

19. A method for preparing a glycoprotein which is a complex or hybrid N-glycan, comprising:

culturing a cell which produces a precursor high-mannose glycoprotein and which contains a heterologous DNA sequence which encodes a protein having the amino acid sequence of formula I.

20. The method of Claim 19, wherein said heterologous DNA sequence has the formula III.

21. The method of Claim 19, wherein said heterologous DNA sequence has the formula IV.

22. A method for preparing a glycoprotein which is a complex or hybrid N-glycan, comprising:

culturing a cell, which produces a precursor high-mannose glycoprotein and which contains a heterologous DNA sequence which encodes a protein having the amino acid sequence of formula II.

23. The method of Claim 22, wherein said heterologous DNA sequence has the formula V.

24. The method of Claim 23, wherein said heterologous DNA sequence has the formula VI.

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Peptide 1:

1                   10                   20                   30  
W A L G Q I F H N F N Y P A A V V V E D D L E V A P D F F E Y f q

Peptide 2:

1                   10  
L W A E L E P K W P K a

Peptide 3:

1                   10  
F W D D W M R R P E Q

Peptide 4:

1  
T D F F P e

Peptide 5:

1                   10  
D L S Y L Q Q E A Y D R D F l

Peptide 6:

1                   10                   20  
L F R G R R V H L A P P O T W D G Y D P S W t

Peptide 7:

1  
L G W L

Peptide 8:

1  
A T Y P L

Oligonucleotides:

2S: 5'-TGG GCI GAA CTI GAA CCI AAA TGG-3'  
      G T           G           G

2A: 5'-CCA TTT IGG TTC IAG TTC IGC CCA-3'  
      C           C           A C

3S: 5'-TTT TGG GAT GAT TGG ATG CG-3'  
      C           C           A

3A: 5'-CG CAT CCA ATC ATC CCA AAA-3'  
      T           G           G

6S: 5'-CAA ACI TGG GAT GGI TAT GAT CC-3'  
      G           C           C           C

6A: 5'-GG ATC ATA ICC ATC CCA IGT TTG-3'  
      G           G           G           C

FIGURE 1

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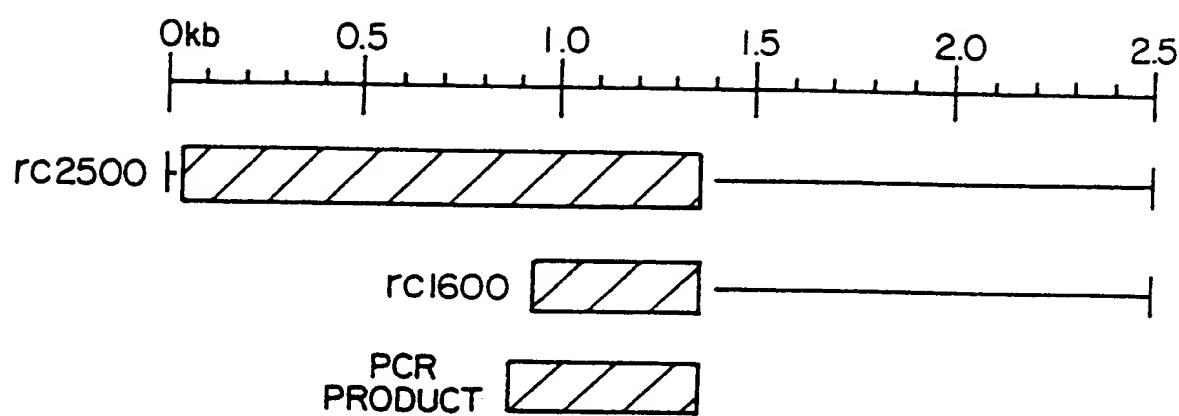


FIGURE 2

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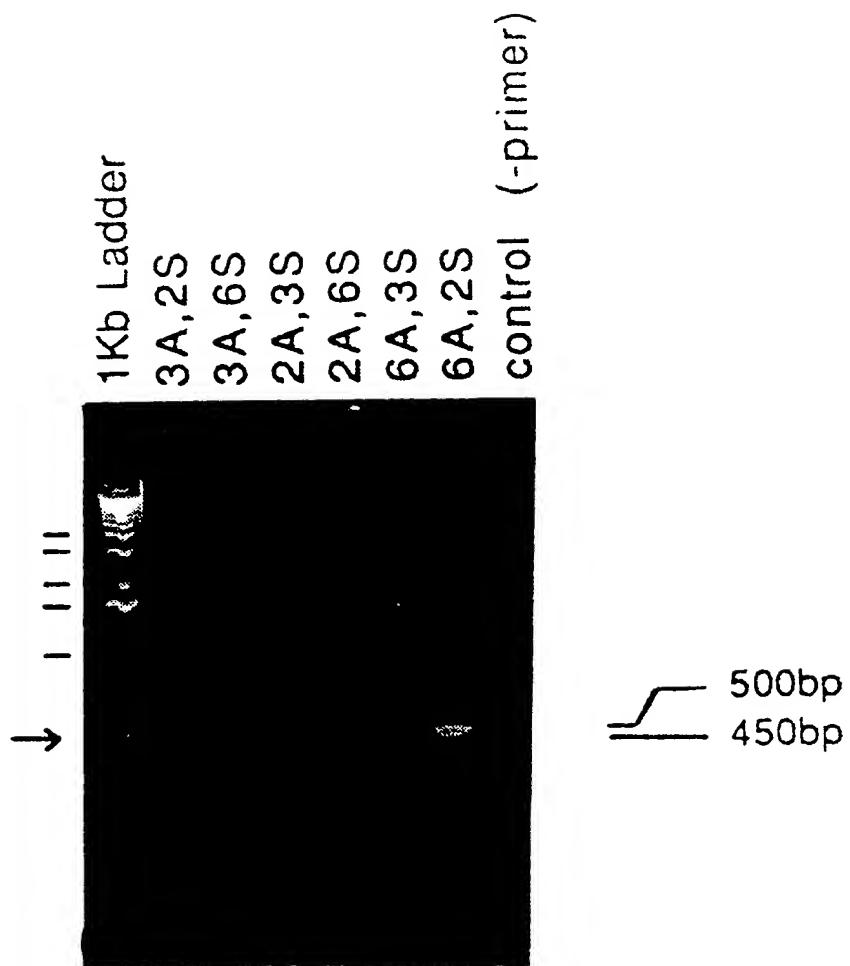


FIGURE 3

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1 gaattccggc aagtcatacc tttgcctgcc ctccccgtg ggggccagg

50: atg ctg aag aag cag tct gct ggg ctt gtg ctg tgg ggt gct atc  
MET LEU LYS LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE

95: ctc ttt gtg gcc tgg aat gcc ctg ctg ctc ctc ttc ttc tgg aca  
LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR

140: cgt cca gtg cct agc agg ctg ccg tca gac aat gct ctc gat gat  
ARG PRO VAL PRO SER ARG LEU PRO SER ASP ASN ALA LEU ASP ASP

185: gac cct gcc agc ctc acc cgt gag gtg atc cgc tta gct cag gat  
ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP

230: gcc gag gta gag ttg gaa cgt cag cgg gga ctg ttg cag cag att  
ALA GLU VAL GLU LEU GLU ARG GLN ARG GLY LEU LEU GLN GLN ILE

275: agg gag cac cat gct ctt tgg agc cag cgg tgg aag gtg cct act  
ARG GLU HIS HIS ALA LEU TRP SER GLN ARG TRP LYS VAL PRO THR

320: gca gcc cct cct gct cag ccg cat gtg cct gtg acc cca ccg cca  
ALA ALA PRO PRO ALA GLN PRO HIS VAL PRO VAL THR PRO PRO PRO

365: gct gtg atc ccc atc ctg gta att gcc tgt gac cgc agc acc gtc  
ALA VAL ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL

410: cgc cgc tgt ttg gac aag cta ctg cat tat cgg cct tca gct gag  
ARG ARG CYS LEU ASP LYS LEU HIS TYR ARG PRO SER ALA GLU

455: ctg ttc ccc atc att gtc agc cag gac tgt ggg cat gag gag aca  
LEU PHE PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR

500: gcc cag gtc att gct tcc tat ggc agc gca gtc aca cac atc cgg  
ALA GLN VAL ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG

545: caa cct gac ctg agc aac att gct gtg cag ccc gac cac cgc aag  
GLN PRO ASP LEU SER ASN ILE ALA VAL GLN PRO ASP HIS ARG LYS

590: ttc cag ggc tac tac aag atc gca cgg cat tac cgc tgg gca ttg  
PHE GLN GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU

635: ggc caa atc ttc cac aat ttc aac tac cca gca gct gtg gtg gtg  
GLY GLN ILE PHE HIS ASN PHE ASN TYR PRO ALA ALA VAL VAL VAL

680: gaa gat gat ctc gag gtg gca cca gac ttc ttt gag tac ttc cag  
GLU ASP ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE GLN

725: gcc act tac cca ctg ttg aaa gca gac ccc tcc ctc tgg tgt gtg  
ALA THR TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL

770: tct gcc tgg aat gac aat ggc aaa gaa cag atg gta gac tcg agt  
SER ALA TRP ASN ASP ASN GLY LYS GLN MET VAL ASP SER SER

815: aag cca gag tta ctc tac cgc aca gat ttc ttt cct ggc tta ggc  
LYS PRO GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY

FIGURE 4

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860: tgg tta ctg ttg gct gaa ctc tgg gct gaa ctg gag ccc aag tgg  
TRP LEU LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP

905: ccc aaa gcc ttc tgg gat gac tgg atg cgc cgg cct gag cag cga  
PRO LYS ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG

950: aag ggg agg gcc tgt gtg cgt cca gaa atc tca aga aca atg aca  
LYS GLY ARG ALA CYS VAL ARG PRO GLU ILE SER ARG THR MET THR

995: ttt ggc cgg aag ggt gtg agc cat ggg cag ttc ttt gac cag cat  
PHE GLY ARG LYS GLY VAL SER HIS GLY GLN PHE PHE ASP GLN HIS

1040: ctc aag ttc atc aag ctg aac cag cag ttt gta ccc ttc acc cag  
LEU LYS PHE ILE LYS LEU ASN GLN GLN PHE VAL PRO PHE THR GLN

1085: ctg gac ctg tcg tac ctt cag cag gag gcc tat gac cgg gat ttc  
LEU ASP LEU SER TYR LEU GLN GLN GLU ALA TYR ASP ARG ASP PHE

1130: ctt gct cgt gtt tat ggt gct ccc cag tta cag gtg gag aaa gtg  
LEU ALA ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL

1175: agg acc aat gac cgg aag gag cta gga gag gtg cgc gta cag tac  
ARG THR ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR

1220: aca ggc agg gac agc ttc aag gct ttc gcc aag gcc ctg ggt gtc  
THR GLY ARG ASP SER PHE LYS ALA PHE ALA LYS ALU GLY VAL

1265: atg gat gac ctc aaa tca ggt gta ccc agg gct gga tac cgg ggc  
MET ASP ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY

1310: att gtc acc ttc tta ttc cgg ggc cgc cgt gtc cac ctg gcg ccc  
ILE VAL THR PHE LEU PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO

1355: cct cag act tgg gat ggc tat gat cct agt tgg act  
PRO GLN THR TRP ASP GLY TYR ASP PRO SER TRP THR

1391 taacagctcc tgcctgtccc ttctgggctc cttcccttgca atttcatgat ctaagatggg  
1451 accgtagtcc ctgggctgca ttgtcttttc tgtctttccc tcttgggtcc atttttttt  
1511 ttttcttttt tgagtggcat ttgaatacac acatgacaag gtgagggttc ttttgtaaa  
1571 ggagtttagat cagggaaagc attctgctgt ctgttggta tcaaggcagca aaccactgtg  
1631 tgatagggga agaatggct ttttggggcc agaaatatcc atgttctgag ttttcttctt  
1691 aggtcatctg cagaggagtt ggcacttta gctttttta ccaggcctt tctttctgac  
1751 ctgagagcca gggcatgaga ctcttggc atgctccctt ttacccccc ctaataaggg  
1811 tctgggctac aggagaatg aacatattgt ggccagaata atactaacca gagggggctc  
1871 attgtcagag tctaggtgca gttattgggt tgtcagagtt aatgccttct gttcttctt  
1931 ctttatctt gacttctgtc agcttttctt tctttgcagc ctagcaattt ttggttctaa  
1991 gataaaaat gaagaggaaa agaaatattt gcacccagct attgggagaa aggtagtgaa  
2051 aaaaaaaactt cattgtacca cttcaaagag acactttga cctttccctt tctaaaaattt  
2111 agtccccctcc ctgttgcctc aggagaatgc tgtgctggc agttctgtgt gatccttctt  
2171 ccctgagttt tatacacagg ctccctcccta aggctgtggc ttctggtgcc cctccctgaca  
2231 taagttacag tggccaagac caggacaact ccggccatga gctaagtctt gcctaccttc  
2291 tccaaaacat tcccatgtcc tcacaggcta ggtatgcagat gttgggttggaa gaggaatttg  
2351 tgtgtgtgtg tgtgtgtgtg tgtgtttct tgcctgaccc cagtttcatg gatgaaaagt  
2411 ggaagctaca gaattatttt caaaaataaa ggctgaattt tctgaaaaaaa aaaaaaaaaa  
2471 aaaaaaccgg aattt

FIGURE 4 (continued)

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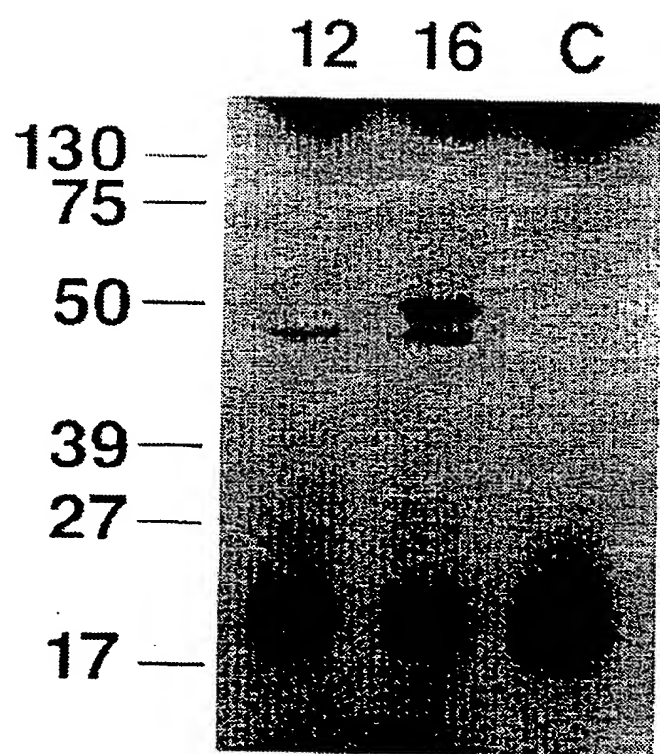


FIGURE 5

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1 aagtttgaa tgaaaaatttattttat tattttctaaa tattttctca tttctctggc  
61 ttttgttaagt agggttttct catccatgtt ttcttcattcat gagttatgg tggatatgaa  
121 ggctatccat tagtataatgt tgatTTTtat attacacttc cttgcgtcagt tcattattga  
181 ttcttttga gtttccagg catattctca caagtaaaga taatagaat agttgcttc  
241 ctcccactt ctgttttga ttttttttc ttggttcatt tgcatggct gcttcccca  
301 gcaaaatgtt aaataaccct ggagatgtt ggcaacttcg tttgtctct gacattcg  
361 ggggcctct ggtgtttccc tttttggtaag gggtaactg tagccctgag gtggacatt  
421 tgatTTTaaa aatcagtcat cttggggcgc tttaggtttaga gaaatggtag gcagatgt  
481 tcactccctg cccctccct cctcccttcc acctggaggg gaaatgaaat ctgacaggta  
541 gaaagagggg agttggggtt cttttctct cttccctccac cagcatcaact ctctgcctct  
601 ccctcaaaaa tacgttctcg ggtcaggata tatgttgcact ccctagagag ctctggagtc  
661 aacccctgg cttccctcca ccctcaactt tggcccccatttcccttacc  
721 tgtggggcat ggagccacga gccccgtgtt gacgggttgc tttctctctc ctgtcttttag  
781 gtgcattggct gccccctaat cccatagtc agaggaggca tcccttaggac tgcgggcaag  
841 ggagccgcaa gcccaggca gcccatttttttgc ctgccttccg gtgggggcca  
901 ggatgtgaa gaagcagtttgc gcaaggcttgc tgctgtgggg cgctatccctc tttgtggcct  
961 ggaatgcctt gctgtcttc ttcttcttggca cgcgcaggc acctggcagg ccaccctcag  
1021 tcagcgctct cgtggcggac cccgcaggcc tcaccggga agtgcatttcg ctggcccaag  
1081 acgcggaggt ggagctggag cgcaggctg ggctgtgc gcaagatcggtt gatgcctgt  
1141 cgagccagcg ggggagggtt gccccccggc cccatcccgcc ccagccgcgt gtgcctgtga  
1201 ccccccggcc ggcgggttatttccatccttgc tcatgccttgc tgaccgcagg actgttgc  
1261 gctgccttggca caagctgttgc cattatcgcc cctcgctga gctttccccc atcatcgta  
1321 gcccaggactg cgggcacggag gagacggccc aggccatcgc ctcctacggc agcgcggta  
1381 cgcacatccg gcagcccgac ctgagcagca ttgcgttgc gccggaccac cgcaagttcc  
1441 agggctacta caagatcgcc cggcactacc gctggcgctt gggccaggc ttccggcagt  
1501 ttccgtttcccg cgcggccgtt gttggggagg atgaccttggca gttggccccc gacttcttgc  
1561 agtactttcg ggccacctat ccgctgttgc agggccaccc cttccctgttgg tgcgttgc  
1621 cctggaaatga caacggcaag gaggcagatgg tggacccagg caggccttgc ctgtcttacc  
1681 gcacccgactt ttcccttggc ctgggttgc tgctgttgc cggactcttgc gctgagctgg  
1741 agcccaagtg gccaaaggcc ttctgggacg actggatgcg gcccggccggag cagcggcagg  
1801 ggcggggctt catabccctt gagatctcaaa gaacgatgcg ctttggccgc aagggtgtga  
1861 cgcacggca gttcttgcg cagcacccatca agttatcaaa gctgaaccagg cagtttgtc  
1921 acttcaccca gctggacccctg ttttacccgc agcggggaggc ctatgaccga gatttccctcg  
1981 cccgcgtcta cgggttccctt cagctgttgc tggagaaagt gaggaccaat gaccggagg  
2041 agctggggga ggtgcgggtt cgtataacgg ggagggacag cttcaaggctt ttgcggccagg  
2101 ctctgggtgt tatggatgc cttaaatgcg gggttcccgag agtggcttac cgggggtatttgc  
2161 tcaccccttcca gttccggggc cggcgttgc acctggcggcc cccaccgcacg tggagggct  
2221 atgatcccttgc ctggaaatttgc cacctgccttgc ttcttcttgc gcccccttgc gccacatca  
2281 gaggttgggtt gaccacatgc cccaggcttgc atcgcccttgc ctgtgtttcc ctcttaggtt  
2341 catttatctt tttgatTTTcc cggatggca tttaatgttca caaatgataa caagaggatt  
2401 atttccctgt tctcaaggaa gtcagatcg gggaaactatt ctgggtatgc ttgcggggta  
2461 ttaaggcagga aaacactgtt tggtgggggg cactggggctt gttggggccca caaatgttcca  
2521 cgtccctgagc tttctcttgc agcatgttgc gagatggggcaacatgttgc tctcttgc  
2581 agaccccttc tccctgactt gctcttcccg ccaggcacga gccccttcc tataactgtt  
2641 ccccttccca gttggggactt agttatggca gaaaggggaca tattttgttgc caaatgata  
2701 ctaacccaaag gggcttccctt gtcaggccctt ggtggagttt gttgggttgc ggggcttact  
2761 gcctcccttgc ttttctcttgc gtcgacccctt cacttagccctt ttctcttgc gcaaggcttgc  
2821 agtttatagt tctgagatgg aaagtttgc ggggcaagca agaccccttgc tcaaggccat  
2881 cccagctgttgc aggagaggagg tgcaggccctt aaggcccttgc gttggggccaa cttcttctt  
2941 gccttacccctt cagagaggac tatggcccttgc ccccttccctt ctgaaaatca gtggccccc  
3001 ttttgccttgc ggaggccctt gttggccctt gttggccctt gttggccctt gttggccctt  
3061 ttttcccttgc gggtttgc gacaggcttgc ttcagcatg aggtggagca gtgaccagg  
3121 ggagcagtta ccaggacccctt gttggccctt gttggccctt gttggccctt gttggccctt  
3181 cgcggccatgtt cttccacaggc caggacccctt gttggccctt gttggccctt gttggccctt

FIGURE 6

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1: MET LEU LYS LYS GLN SER ALA GLY LEU VAL LEU TRP GLY ALA ILE  
16: LEU PHE VAL ALA TRP ASN ALA LEU LEU LEU PHE PHE TRP THR  
31: ARG PRO ALA PRO GLY ARG PRO PRO SER VAL SER ALA LEU ASP GLY  
46: ASP PRO ALA SER LEU THR ARG GLU VAL ILE ARG LEU ALA GLN ASP  
61: ALA GLU VAL GLU LEU GLU ARG ARG GLY LEU LEU GLN GLN ILE  
76: GLY ASP ALA LEU SER SER GLN ARG GLY ARG VAL PRO THR ALA ALA  
91: PRO PRO ALA GLN PRO ARG VAL PRO VAL THR PRO ALA PRO ALA VAL  
106: ILE PRO ILE LEU VAL ILE ALA CYS ASP ARG SER THR VAL ARG ARG  
121: CYS LEU ASP LYS LEU LEU HIS TYR ARG PRO SER ALA GLU LEU PHE  
136: PRO ILE ILE VAL SER GLN ASP CYS GLY HIS GLU GLU THR ALA GLN  
151: ALA ILE ALA SER TYR GLY SER ALA VAL THR HIS ILE ARG GLN PRO  
166: ASP LEU SER SER ILE ALA VAL PRO PRO ASP HIS ARG LYS PHE GLN  
181: GLY TYR TYR LYS ILE ALA ARG HIS TYR ARG TRP ALA LEU GLY GLN  
196: VAL PHE ARG GLN PHE ARG PHE PRO ALA ALA VAL VAL VAL GLU ASP  
211: ASP LEU GLU VAL ALA PRO ASP PHE PHE GLU TYR PHE ARG ALA THR  
226: TYR PRO LEU LEU LYS ALA ASP PRO SER LEU TRP CYS VAL SER ALA  
241: TRP ASN ASP ASN GLY LYS GLU GLN MET VAL ASP ALA SER ARG PRO  
256: GLU LEU LEU TYR ARG THR ASP PHE PHE PRO GLY LEU GLY TRP LEU  
271: LEU LEU ALA GLU LEU TRP ALA GLU LEU GLU PRO LYS TRP PRO LYS  
286: ALA PHE TRP ASP ASP TRP MET ARG ARG PRO GLU GLN ARG GLN GLY  
301: ARG ALA CYS ILE ARG PRO GLU ILE SER ARG THR MET THR PHE GLY  
316: ARG LYS GLY VAL THR HIS GLY GLN PHE PHE ASP GLN HIS LEU LYS  
331: PHE ILE LYS LEU ASN GLN GLN PHE VAL HIS PHE THR GLN LEU ASP  
346: LEU SER TYR LEU GLN ARG GLU ALA TYR ASP ARG ASP PHE LEU ALA  
361: ARG VAL TYR GLY ALA PRO GLN LEU GLN VAL GLU LYS VAL ARG THR  
376: ASN ASP ARG LYS GLU LEU GLY GLU VAL ARG VAL GLN TYR THR GLY  
391: ARG ASP SER PHE LYS ALA PHE ALA LYS ALA LEU GLY VAL MET ASP  
406: ASP LEU LYS SER GLY VAL PRO ARG ALA GLY TYR ARG GLY ILE VAL  
421: THR PHE GLN PHE ARG GLY ARG ARG VAL HIS LEU ALA PRO PRO PRO  
436: THR TRP GLU GLY TYR ASP PRO SER TRP ASN\*\*\*

FIGURE 7

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## START

880: c c tgc cct ccg gtg ggg gcc agg|atg ctg aag aag cag tct gca  
 3: . . CYS PRO PRO VAL GLY ALA ARG MET LEU LYS LYS GLN SER ALA

924: ggg ctt gtg ctg tgg ggc gct atc ctc ttt gtg gcc tgg aat gcc  
 3: GLY LEU VAL LEU TRP GLY ALA ILE LEU PHE VAL ALA TRP ASN ALA

969: ctg ctg ctc ctc ttc tgg acg cgc cca gca cct ggc agg cca  
 3: LEU LEU LEU LEU PHE PHE TRP THR ARG PRO ALA PRO GLY ARG PRO

1014: ccc tca gtc agc gct ctc gat ggc gac ccc gcc agc ctc acc cgg  
 3: PRO SER VAL SER ALA LEU ASP GLY ASP PRO ALA SER LEU THR ARG

1059: gaa gtg att cgc ctg gcc caa gac gcc gag gtg gag ctg gag cgc  
 3: GLU VAL ILE ARG LEU ALA GLN ASP ALA GLU VAL GLU LEU GLU ARG

1104: agg cgt ggg ctg ctg cag cag atc ggg gat gcc ctg tcg agc cag  
 3: ARG ARG GLY LEU LEU GLN GLN ILE GLY ASP ALA LEU SER SER GLN

1149: cgg ggg agg gtg ccc acc gcg gcc cct ccc gcc cag cgc cgt gtg  
 3: ARG GLY ARG VAL PRO THR ALA ALA PRO PRO ALA GLN PRO ARG VAL

1194: cct gtg acc ccc gcg ccg gcg gtg att ccc atc ctg gtc atc gcc  
 3: PRO VAL THR PRO ALA PRO ALA VAL ILE PRO ILE LEU VAL ILE ALA

1239: tgt gac cgc agc act gtt cgg cgc tgc ctg gac aag ctg ctg cat  
 3: CYS ASP ARG SER THR VAL ARG ARG CYS LEU ASP LYS LEU LEU HIS

1284: tat cgg ccc tcg gct gag ctc ttc ccc atc atc gtt agc cag gac  
 3: TYR ARG PRO SER ALA GLU LEU PHE PRO ILE ILE VAL SER GLN ASP

1329: tgc ggg cac gag gag acg gcc cag gcc atc gcc tcc tac ggc agc  
 3: CYS GLY HIS GLU GLU THR ALA GLN ALA ILE ALA SER TYR GLY SER

1374: gcg gtc acg cac atc cgg cag ccc gac ctg agc agc att gcg gtg  
 3: ALA VAL THR HIS ILE ARG GLN PRO ASP LEU SER SER ILE ALA VAL

1419: ccg ccg gac cac cgc aag ttc cag ggc tac tac aag atc gcg cgc  
 3: PRO PRO ASP HIS ARG LYS PHE GLN GLY TYR TYR LYS ILE ALA ARG

1464: cac tac cgc tgg gcg ctg ggc cag gtc ttc cgg cag ttt cgc ttc  
 3: HIS TYR ARG TRP ALA LEU GLY GLN VAL PHE ARG GLN PHE ARG PHE

1509: ccc gcg gcc gtg gtg gtg gag gat gac ctg gag gtg gcc ccc gac  
 3: PRO ALA ALA VAL VAL VAL GLU ASP ASP LEU GLU VAL ALA PRO ASP

1554: ttc ttc gag tac ttt cgg gcc acc tat ccg ctg ctg aag gcc gac  
 3: PHE PHE GLU TYR PHE ARG ALA THR TYR PRO LEU LEU LYS ALA ASP

1599: ccc tcc ctg tgg tgc tgc tgg aat gac aac ggc aag gag  
 3: PRO SER LEU TRP CYS VAL SER ALA TRP ASN ASP ASN GLY LYS GLU

1644: cag atg gtg gac gcc agc agg cct gag ctg ctc tac cgc acc gac  
 3: GLN MET VAL ASP ALA SER ARG PRO GLU LEU LEU TYR ARG THR ASP

FIGURE 8

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1689: ttt ttc cct ggc ctg ggc tgg ctg ctg ttg gcc gag ctc tgg gct  
3: PHE PHE PRO GLY LEU GLY TRP LEU LEU LEU ALA GLU LEU TRP ALA

1734: gag ctg gag ccc aag tgg cca aag gcc ttc tgg gac gac tgg atg  
3: GLU LEU GLU PRO LYS TRP PRO LYS ALA PHE TRP ASP ASP TRP MET

1779: cgg cgg ccg gag cag cgg cag ggg cgg gcc tgc ata cgc cct gag  
3: ARG ARG PRO GLU GLN ARG GLN GLY ARG ALA CYS ILE ARG PRO GLU

1824: atc tca aga acg atg acc ttt ggc cgc aag ggt gtg acg cac ggg  
3: ILE SER ARG THR MET THR PHE GLY ARG LYS GLY VAL THR HIS GLY

1869: cag ttc ttt gac cag cac ctc aag ttt atc aag ctg aac cag cag  
3: GLN PHE PHE ASP GLN HIS LEU LYS PHE ILE LYS LEU ASN GLN GLN

1914: ttt gtg cac ttc acc cag ctg gac ctg tct tac ctg cag cgg gag  
3: PHE VAL HIS PHE THR GLN LEU ASP LEU SER TYR LEU GLN ARG GLU

1959: gcc tat gac cga gat ttc ctc gcc cgc gtc tac ggt gct ccc cag  
3: ALA TYR ASP ARG ASP PHE LEU ALA ARG VAL TYR GLY ALA PRO GLN

2004: ctg cag gtg gag aaa gtg agg acc aat gac cgg aag gag ctg ggg  
3: LEU GLN VAL GLU LYS VAL ARG THR ASN ASP ARG LYS GLU LEU GLY

2049: gag gtg cgg gtg cag tat acg ggg agg gac agc ttc aag gct ttc  
3: GLU VAL ARG VAL GLN TYR THR GLY ARG ASP SER PHE LYS ALA PHE

2094: gcc aag gct ctg ggt gtt atg gat gac ctt aag tcg ggg gtt ccg  
3: ALA LYS ALA LEU GLY VAL MET ASP ASP LEU LYS SER GLY VAL PRO

2139: aga gct ggc tac cgg ggt att gtc acc ttc cag tcc cgg ggc cgc  
3: ARG ALA GLY TYR ARG GLY ILE VAL THR PHE GLN PHE ARG GLY ARG

2184: cgt gtc cac ctg gcg ccc cca ccg acg tgg gag ggc tat gat cct  
3: ARG VAL HIS LEU ALA PRO PRO PRO THR TRP GLU GLY TYR ASP PRO  
STOP

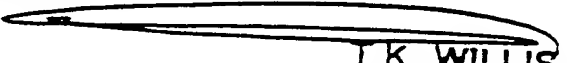
2229: agc tgg aat | tag cac ctg cct g  
3: SER TRP ASN \*\*\* HIS LEU PRO .

FIGURE 8 (continued)

## SUBSTITUTE SHEET

# INTERNATIONAL SEARCH REPORT

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<b>I. CLASSIFICATION OF SUBJECT MATTER</b> (if several classification symbols apply, indicate all)*		
According to International Patent Classification (IPC) or to both National Classification and IPC Int.C1.5 C 12 N 15/54 C 12 N 9/10		
<b>II. FIELDS SEARCHED</b>		
Minimum Documentation Searched <sup>1</sup>		
Classification System	Classification Symbols	
Int.C1.5	C 12 N	
Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched <sup>2</sup>		
<b>III. DOCUMENTS CONSIDERED TO BE RELEVANT*</b>		
Category <sup>3</sup>	Citation of Document <sup>4</sup> , <sup>11</sup> with indication, where appropriate, of the relevant passages <sup>12</sup>	Relevant to Claim No. <sup>13</sup>
X	Glycoconjugate Journal, vol. 7, no. 5, 10 October 1990, (Lund, SE), E. HULL et al.: "Isolation of 13 and 15 kilobase human genomic DNA clones containing the gene for UDP-N-acetylglucosamine:alpha-3-D-mannoside beta-1,2-N-acetylglucosaminyltransferase I", page 468, abstract no. 85, see the whole document ---	1-3,7-9 ,13-15
X	Glycoconjugate Journal, vol. 7, no. 5, 10 October 1990, (Lund, SE), M. SARKAR et al.: "Rabbit liver UDP-N-acetylglucosamine:alpha-3-D-mannoside beta-1,2-N-acetylglucosaminyltransferase I: characterization of a 2,5 kilobase cDNA clone", page 380, abstract no. 4, see the whole document ---	1-3,7-9 ,13-15
* Special categories of cited documents : <sup>10</sup> "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reasons (as specified) "O" document referring to an oral disclosure, etc, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed		"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "Z" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "A" document member of the same patent family
<b>IV. CERTIFICATION</b>		
Date of the Actual Completion of the International Search	Date of Mailing of this International Search Report	
18-02-1992	15.04.93	
International Searching Authority <b>EUROPEAN PATENT OFFICE</b>	Signature of Authorized Officer  T.K. WILLIS	

## INTERNATIONAL SEARCH REPORT

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## III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)

Category	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No.
P, X	<p>Proc. Natl. Acad. Sci. USA, vol. 88, no. 1, January 1991, Natl. Acad. Sci., (Washington, DC, US), M. SAKKAR et al.: "Molecular cloning and expression of cDNA encoding the enzyme that controls conversion of high-mannose to hybrid and complex N-glycans: UDP-N-acetylglucosamine:alpha-3-D-mannoside beta-1,2-N-acetylglucosaminyltransferase I", pages 234-238, see figure 4; page 236, left-hand column, line 26 - page 237, right-hand column, line 6</p> <p>---</p>	1-3,7-9 .13-15
P, X	<p>Biochem. Soc. Trans., vol. 19, no. 3, August 1991, Biochemical Society, (London, GB), H. SCHACHTER et al.: "Molecular cloning of human and rabbit UDP-N-acetylglucosamine:alpha-3-D-mannoside beta-1,2-N-acetylglucosaminyltransferase I", pages 645-648, see page 646, left-hand column, line 1 - page 648, right-hand column, line 23</p> <p>---</p>	1-3,7-9 .13-15
Y	<p>J. Biol. Chem., vol. 263, no. 17, 15 June 1988, Am. Soc. Biol. Chem., Inc., (US), Y. NISHIKAWA et al.: "Control of glycoprotein synthesis. Purification and characterization of rabbit liver UDP-N-acetylglucosamine:alpha-3-D-mannoside beta-1,2-N-acetylglucosaminyltransferase I", pages 8270-8281, see table I; abstract; page 8270, right-hand column, lines 25-29 (cited in the application)</p> <p>---</p>	1-3,7-9 .13-15, 19-21
Y	<p>J. Biol. Chem., vol. 265, no. 2, 15 January 1990, Am. Soc. Biol. Chem., Inc., (US), F. YAMAMOTO et al.: "Cloning and characterization of DNA complementary to human UDP-GalNAc: Fucalph1 2Gal alph1 3GalNAc transferase (histo-blood group A transferase) mRNA", pages 1146-1151, see materials and methods (cited in the application)</p> <p>---</p>	1-3,7-9 .13-15, 19-21 --

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III. DOCUMENTS CONSIDERED TO BE RELEVANT		(CONTINUED FROM THE SECOND SHEET)
Category *	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No.
Y	J. Biol. Chem., vol. 256, no. 2, 25 January 1981, Am. Soc. Biol. Chem., Inc., (US), C.L. OPPENHEIMER et al.: "Purification and characterization of a rabbit liver alphal 3 mannoside betal 2 N-acetylglucosaminyltransferase", pages 799-804, see page 801, left-hand column, line 8 - right-hand column, line 8 (cited in the application) -----	1-3,7-9 ,13-15, 19-21